# PCT

(22) International Filing Date:

# WORLD INTELLECTUAL PROPERTY ORGANIZATION International Bureau



# INTERNATIONAL APPLICATION PUBLISHED UNDER THE PATENT COOPERATION TREATY (PCT

	(51) International Patent Classification 6;		TATELY COOPERATION TREATY (PCT)
		A1	(11) International Publication Number: WO 96/37609
i	C12N 15/12, 15/85, 15/62, C07K 14/72, 19/00, C12N 5/10, A61K 38/16		(43) International Publication Date: 28 November 1996 (28.11.96)
	(21) International Application Number: PCT/GBS	96/0119	95 (81) Designated States: AL AM AT ALL AZ DR BC DR DE

20 May 1996 (20.05.96)

(30) Priority Data:
9510759.5
26 May 1995 (26.05.95)
9513882.2
7 July 1995 (07.07.95)
9517316.7
24 August 1995 (24.08.95)
9605656.9
18 March 1996 (18.03.96)
GB

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81) Designated States: AL, AM, AT, AU, AZ, BB, BG, BR, BY, CA, CH, CN, CZ, DE, DK, EE, ES, FI, GB, GE, HU, IS, JP, KE, KG, KP, KR, KZ, LK, LR, LS, LT, LU, LV, MD, MG, MK, MN, MW, MX, NO, NZ, PL, PT, RO, RU, SD, SE, SG, SI, SK, TJ, TM, TR, TT, UA, UG, US, UZ, VN, ARIPO patent (KE, LS, MW, SD, SZ, UG), Eurasian patent (AM, AZ, BY, KG, KZ, MD, RU, TJ, TM), European patent (AT, BE, CH, DE, DK, ES, FI, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE), OAPI patent (BF, BJ, CF, CG, CI, CM, GA, GN, ML, MR, NE, SN, TD, TG).

Published

With international search report.

(54) Title: A GENE SWITCH COMPRISING AN ECDYSONE RECEPTOR

(57) Abstract

The invention relates to an insect steroid receptor protein which is capable of acting as a gene switch which is responsive to a chemical inducer enabling external control of the gene.

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# A gene switch comprising an ecdysone receptor

The present inventi n relates to the identification and characterization of insect steroid receptors from the Lepidoptera species Heliothis virescens, and the nucleic acid encoding therefor. The present invention also relates to the use of such receptors, and such nucleic acid, particularly, but not exclusively, in screening methods, and gene switches.

By gene switch we mean a gene sequence which is responsive to an applied exogenous chemical inducer enabling external control of expression of the gene controlled by said gene sequence.

Lipophilic hormones such as steroids induce changes in gene expression to elicit profound effects on growth, cellular differentiation, and homeostasis. These hormones recognise intracellular receptors that share a common modular structure consisting of three main functional domains: a variable amino terminal region that contains a transactivation domain, a DNA binding domain, and a ligand binding domain on the carboxyl side of the molecule. The DNA binding domain contains nine invariant cysteines, eight of which are involved in zinc coordination to form a two-finger structure. In the nucleus the hormone-receptor complex binds to specific enhancer-like sequences called hormone response elements (HREs) to modulate transcription of target genes.

The field of insect steroid research has undergone a revolution in the last three years as a result of the cloning and preliminary characterisation of the first steroid receptor member genes. These developments suggest the time is ripe to try to use this knowledge to improve our tools in the constant fight against insect pests. Most of the research carried out on the molecular biology of the steroid receptor superfamily has been on *Drosophila melanogaster* (Diptera), see for example International Patent Publication No WO91/13167, with some in *Manduca* and *Galleria* (Lepidoptera).

It has been three decades since 20-hydroxyecdysone was first isolated and shown to be involved in the regulation of development of insects. Since then work has been carried out to try to understand the pathway by which this small hydrophobic molecule regulates a number of activities. By the early 1970s, through the studies of Clever and Ashburner, it was clear that at least in the salivary glands of third instar *Drosophila* larvae, the application of ecdysone lead to the reproducible activation of over a hundred genes. The ecdysone receptor in this pathway is involved in the regulation of two classes of genes: a small class (early genes) which are induced by the ecdysone receptor and a large class (late genes) which are repressed by the ecdysone receptor. The early class of genes are thought to have two functions reciprocal to those of the ecdysone receptor; the repression of the early transcripts and the induction of late gene transcription. Members of the early genes so far isolated and characterised belong to the class of molecules with characteristics similar to known

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transcription factors. They are thus predicted to behave as expected by the model f ecdysone action (Ashburner, 1991). More recently, the early genes E74 and E75 have been shown to bind both types of ecdysone inducible genes (Thurnmel et al., 1990; Segraves and Hogness, 1991), thus supporting their proposed dual activities. It should be noted however, that the activation of a hierarchy of genes is not limited to third instar larvae salivary glands, but that the response to the ecdysone peak at the end of larval life is observed in many other tissues, such as the imaginal disks (i.e. those tissues which metamorphose to adult structures) and other larval tissues which histolyse at the end of larval life (eg. larval fat body). The model for ecdysone action as deduced by studying the third instar chromosome puffing may not apply to the activation of ecdysone regulated genes in adults. In other words, the requirement for other factors in addition to the active ecdysone receptor must be satisfied for correct developmental expression (e.g. the *Drosophila* yolk protein gene expression in adults is under control of doublesex, the last gene in the sex determination gene hierarchy).

The ecdysone receptor and the early gene E75 belong to the steroid receptor superfamily. Other Drosophila genes, including ultraspiracle, tailless, sevenup and FIZ-FI, also belong to this family. However, of all these genes only the ecdysone receptor is known to have a ligand, and thus the others are known as orphan receptors. Interestingly, despite the ultraspiracle protein ligand binding region sharing 49% identity with the vertebrate retinoic X receptor (RXR) ligand binding region (Oro et al., 1990), they do not share the same ligand (i.e. the RXR ligand is 9-cis retinoic acid) (Heymann et al., 1992 and Mangelsdorf et al., 1992). All the Drosophila genes mentioned are involved in development, ultraspiracle for example, is required for embryonic and larval abdominal development. The protein products of these genes all fit the main features of the steroid receptor superfamily (Evans, 1988; Green and Chambon, 1988, Beato, 1989) i.e. they have a variable N terminus region involved in ligand independent transactivation (Domains A and B), a highly conserved 66-68 amino acid region which is responsible for the binding of DNA at specific sites (Domain C), a hinge region thought to contain a nuclear translocation signal (Domain D), and a well conserved region containing the ligand binding region, transactivation sequences and the dimerisation phase (Domain E). The last region, domain F, is also very variable and its function is unknown.

Steroid receptor action has been elucidated in considerable detail in vertebrate systems at both the cellular and molecular levels. In the absence of ligand, the receptor molecule resides in the cytoplasm where it is bound by Hsp90, Hsp70, and p59 to form the inactive complex (Evans, 1988). Upon binding of the ligand molecule by the receptor a conformational change takes place which releases the Hsp90, Hsp70 and p59 molecules, while exposing the nuclear translocation signals in the receptor. The ligand dependent conformational change is seen in the ligand binding domain of both progesterone and retinoic acid receptors (Allan et

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al., 1992a). This conformational change has been further characterised in the progesterone receptor and was found to be indispensable for gene transactivation (Allan et al., 1992b). Once inside the nucleus the receptor dimer binds to the receptor responsive element at a specific site on the DNA resulting in the activation or repression of a target gene. The receptor responsive elements usually consist of degenerate direct repeats, with a spacer between 1 and 5 nucleotides, which are bound by a receptor dimer through the DNA binding region (Domain C).

Whereas some steroid hormone receptors are active as homodimers others act as heterodimers. For example, in vertebrates, the retinoic acid receptor (RAR) forms heterodimers with the retinoic X receptor (RXR). RXR can also form heterodimers with the thyroid receptor, vitamin D receptor (Yu et al., 1991; Leid et al., 1992) and peroxisome activator receptor (Kliewer et al., 1992). Functionally the main difference between homodimers and heterodimers is increased specificity of binding to specific response elements. This indicates that different pathways can be linked, co-ordinated and modulated, and more importantly this observation begins to explain the molecular basis of the pleotropic activity of retinoic acid in vertebrate development (Leid et al., 1992b). Similarly, the Drosophila ultraspiracle gene product was recently shown to be capable of forming heterodimers with retinoic acid, thyroid, vitamin D and peroxisome activator receptors and to stimulate the binding of these receptors to their target responsive elements (Yao et al., 1993). More significantly, the ultraspiracle gene product has also been shown to form heterodimers with the ecdysone receptor, resulting in cooperative binding to the ecdysone response element and capable of rendering mammalian cells ecdysone responsive (Yao et al., 1992). The latter is of importance since transactivation of the ecdysone gene alone in mammalian cells fails to elicit an ecdysone response (Koelle et al., 1991), therefore suggesting that the ultraspiracle gene product is an integral component of a functional ecdysone receptor (Yao et al., 1992). It is possible that the ultraspiracle product competes with other steroid receptors or factors to form heterodimers with the ecclysone receptor. Moreover it remains to be investigated if ultraspiracle is expressed in all tissues of the Drosophila larvae. Despite ultraspiracle being necessary to produce a functional ecdysone receptor, the mechanism by which this activation takes place is as yet undetermined.

We have now isolated and characterised the ecdysone steroid receptor from *Heliothis virescens* (hereinafter HEcR). We have found that surprisingly unlike the *Drosophila* ecdysone steroid receptor (hereinafter DEcR), in reports to-date, HEcR can be induced by known non-steroidal inducers. It will be appreciated that this provides many advantages for the system.

Steroids are difficult and expensive to make. In addition, the use of a non-steroid as the inducer allows the system to be used in agrochemical and pharmaceutical applications, not

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least because it avoids application of a steroid which is already present in insects and/or mammals. For example, it would not be feasible to use a gene switch in a mammalian cell which was induced by a naturally occurring steroidal inducer. It will also be appreciated that for environmental reasons it is advantageous to avoid the use of steroids as inducers.

According to one aspect of the present invention there is provided DNA having the sequence shown in Seq ID No. 2, wherein Seq ID No 2 gives the sequence for the HEcR.

According to another aspect of the present invention there is provided DNA having part of the sequence shown in Seq ID No. 2, which encodes for the HEcR ligand binding domain.

According to another aspect of the present invention there is provided DNA having part of the sequence shown in Seq ID No. 2, which encodes for the HECR DNA binding domain.

According to yet another aspect of the present invention there is provided DNA having part of the sequence shown in Seq ID No. 2, which encodes for the HECR transactivation domain.

According to a further aspect of the present invention there is provided DNA having part of the sequence shown in Seq ID No. 2, which encodes for the HEcR hinge domain.

According to a still further aspect of the present invention there is provided DNA having part of the sequence shown in Seq ID No. 2, which encodes for the HECR carboxy terminal region.

According to one aspect of the present invention there is provided DNA having the sequence shown in Seq ID No. 3, wherein Seq ID No 3 gives the sequence for the HEcR.

According to another aspect of the present invention there is provided DNA having part of the sequence shown in Seq ID No. 3, which encodes for the HEcR ligand binding domain.

According to another aspect of the present invention there is provided DNA having part of the sequence shown in Seq ID No. 3, which encodes for the HEcR DNA binding domain.

According to yet another aspect of the present invention there is provided DNA having part of the sequence shown in Seq ID No. 3, which encodes for the HECR transactivation domain.

According to a further aspect of the present invention there is provided DNA having part of the sequence shown in Seq ID No. 3, which encodes for the HEcR hinge domain.

According to a still further aspect of the present invention there is provided DNA having part of the sequence shown in Seq ID No. 3, which encodes for the HEcR carboxy terminal region.

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According to one aspect of the present invention there is provided DNA having the sequence shown in Seq ID No. 4, wherein Seq ID No 4 gives the sequence for the HEcR.

According to another aspect of the present invention there is provided DNA having part of the sequence shown in Seq ID No. 4, which encodes for the HEcR ligand binding domain.

According to another aspect of the present invention there is provided DNA having part of the sequence shown in Seq ID No. 4, which encodes for the HEcR DNA binding domain.

According to yet another aspect of the present invention there is provided DNA having part of the sequence shown in Seq ID No. 4, which encodes for the HEcR transactivation domain.

According to a further aspect of the present invention there is provided DNA having part of the sequence shown in Seq ID No. 4, which encodes for the HEcR hinge domain.

According to a still further aspect of the present invention there is provided DNA having part of the sequence shown in Seq ID No. 4, which encodes for the HEcR carboxy terminal region.

As mentioned above, steroid receptors are eukaryotic transcriptional regulatory factors which, in response to the binding of the steroid hormone, are believed to bind to specific DNA elements and activate transcription. The steroid receptor can be divided into six regions, designated A to F, using alignment techniques based on shared homology with other members of the steroid hormone receptor superfamily. Krust et al identified two main regions in the receptor, C and E. Region C is hydrophilic and is unusual in its high content in cysteine, lysine and arginine. It corresponds to a DNA-binding domain, sometimes referred to as the "zinc finger". It is the DNA binding domain which binds to the upstream DNA of the responsive gene. Such upstream DNA is known as the hormone response element or HRE for short. Region E is hydrophobic and is identified as the hormone (or ligand) binding domain. Region E can be further subdivided into regions E1, E2 and E3.

The region D, which separates domains C and E is highly hydrophobic and is flexible. It is believe that communication between domains E and C involves direct contact between them through region D, which provides a hinge between the two domains. Region D is therefore referred to as the hinge domain.

The mechanism of the receptor appears to require it to interact with some element(s) of the transcription machinery over and above its interactions with the hormone and the hormone response element. N-terminal regions A and B perform such a function and are jointly known as the transactivation domain. The carboxy terminal region is designated F.

The domain boundaries of the HEcR can be defined as follows:

DOMAIN	INTERVALS			
	base pairs	amino acids		
Transactivating (A/B)	114-600	1-162		
DNA Binding (C)	601-798	163-228		
Hinge (D)	· 799-1091	229-326		
Ligand Binding (E)	1092-1757	327-545		
C-Terminal End (F)	1758-1844	546-577		

The DNA binding domain is very well defined and is 66 amino acids long, thus providing good boundaries. The above intervals have been defined using the multiple alignment for the ecdysone receptors (Figure 5).

The present invention also includes DNA which shows homology to the sequences of the present invention. Typically homology is shown when 60% or more of the nucletides are common, more typically 65%, preferably 70%, more preferably 75%, even more preferably 80% or 85%, especially preferred are 90%, 95%, 98% or 99% or more homology.

The present invention also includes DNA which hybridises to the DNA of the present invention and which codes for at least part of the *Heliothis* ecdysone receptor transactivation domain, DNA binding domain, hinge domain, ligand binding domain and/or carboxy terminal region. Preferably such hybridisation occurs at, or between, low and high stringency conditions. In general terms, low stringency conditions can be defined as 3 x SCC at about ambient temperature to about 65°C, and high stringency conditions as 0.1 x SSC at about 65°C. SSC is the name of a buffer of 0.15M NaCl, 0.015M trisodium citrate. 3 x SSC is three time as strong as SSC and so on.

The present invention further includes DNA which is degenerate as a result of the genetic code to the DNA of the present invention and which codes for a polypeptide which is at least part of the *Heliothis* ecdysone receptor transactivation domain, DNA binding domain, hinge domain, ligand binding domain and/or carboxy terminal region.

The DNA of the present invention may be cDNA or DNA which is in an isolated form.

According to another aspect of the present invention there is provided a polypeptide comprising the *Heliothis* ecdysone receptor or a fragment thereof, wherein said polypeptide is substantially free from other proteins with which it is ordinarily associated, and which is coded for by any of the DNA of the present invention.

According to another aspect of the present invention there is provided a polypeptide which has the amino acid sequence of Seq ID No. 4 or any allelic variant or derivative thereof, wherein Seq ID No. 4 gives the amino acid sequence of the HEcR polypeptide.

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According to another aspect of the present invention there is provided a polypeptide which has part of the amino acid sequence of Seq ID No. 4 or any allelic variant or derivative thereof, which sequence provides the HEcR ligand binding domain.

According to another aspect of the present invention there is provided a polypeptide which has part of the amino acid sequence of Seq ID No. 4 or any allelic variant or derivative thereof, which sequence provides the HEcR DNA binding domain.

According to yet another aspect of the present invention there is provided a polypeptide which has part of the amino acid sequence of Seq ID No. 4 or any allelic variant or derivative thereof, which sequence provides the HEcR transactivation domain.

According to a further aspect of the present invention there is provided a polypeptide which has the amino acid sequence of a part of Seq ID No. 4 or any allelic variant or derivative thereof, which sequence provides the HEcR hinge domain.

According to a still further aspect of the present invention there is provided a polypeptide which has the amino acid sequence of a part of Seq ID No. 4 or any allelic variant or derivative thereof, which sequence provides the HEcR carboxy terminal region.

For the avoidance of doubt, spliced variants of the amino acid sequences of the present invention are included in the present invention.

Preferably, said derivative is a homologous variant which has conservative amino acid changes. By conservation amino acid changes we mean replacing an amino acid from one of the amino acid groups, namely hydrophobic, polar, acidic or basic, with an amino acid from within the same group. An examples of such a change is the replacement of valine by methionine and vice versa.

According to another aspect of the present invention there is provided a fusion polypeptide comprising at least one of the polypeptides of the present invention functionally linked to an appropriate non-Heliothis ecdysone receptor domain(s).

According to an especially preferred embodiment of the present invention the HEcR ligand binding domain of the present invention is fused to a DNA binding domain and a transactivation domain.

According to another embodiment of the present invention the DNA binding domain is fused to a ligand binding domain and a transactivation domain.

According to yet another embodiment of the present invention the transactivation domain is fused to a ligand binding domain and a DNA binding domain.

The present invention also provides recombinant DNA encoding for these fused polypeptides.

According to an especially preferred embodiment of the present invention there is provided recombinant nucleic acid comprising a DNA sequence encoding the HECR ligand

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binding domain functionally linked to DNA encoding the DNA binding domain and transactivation domain from a glucocorticoid receptor.

According to yet another aspect of the present invention there is provided recombinant nucleic acid comprising a DNA sequence comprising a reporter gene operably linked to a promoter sequence and a hormone response element which hormone response element is responsive to the DNA bonding domain encoded by the DNA of of the present invention.

According to another aspect of the present invention there is provided a construct transformed with nucleic acid, recombinant DNA, a polypeptide or a fusion polypeptide of the present invention. Such constructs include plasmids and phages suitable for transforming a cell of interest. Such constructs will be well known to those skilled in the art.

According to another aspect of the present invention there is provided a cell transformed with nucleic acid, recombinant DNA, a polypeptide, or a fusion polypeptide of the present invention.

Preferably the cell is a plant, fungus or mammalian cell.

For the avoidance of doubt fungus includes yeast.

The present invention therefore provides a gene switch which is operably linked to a foreign gene or a series of foreign genes whereby expression of said foreign gene or said series of foreign genes may be controlled by application of an effective exogenous inducer.

Analogs of ecdysone, such as Muristerone A, are found in plants and disrupt the development of insects. It is therefore proposed that the receptor of the present invention can be used be in plants transformed therewith as an insect control mechanism. The production of the insect-damaging product being controlled by an exogenous inducer. The insect-damagin g product can be ecdysone or another suitable protein.

The first non-steroidal ecdysteroid agonists, dibenzoyl hydrazines, typified by RH-5849 [1,2-dibenzoyl, 1-tert-butyl hydrazide], which is commercially available as an insecticide from Rohm and Haas, were described back in 1988. Another commercially available compound in this series is RH-5992 [tebufenozide, 3,5-dimethylbenzoic acid 1-1 (1,1-dimethylethyl)-2(4-ethylbenzoyl) hydrazide]. These compounds mimic 20-hydroxyecdysone (20E) in both Manduca sexta and Drosophila melanogaster. These compounds have the advantage that they have the potential to control insects using ecdysteroid agonists which are non-steroidal. Further Examples of such dibenzoyl hydrazines are given in US Patent No. 5,117,057 to Rohm and Haas, and Oikawa et al, Pestic Sci, 41, 139-148 (1994). However, it will be appreciated that any inducer of the gene switch of the present invention, whether steroidal or non-steroidal, and which is currently or becomes available, may be used.

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The gene switch of the present invention, then, when linked to an exogenous or foreign gene and introduced into a plant by transformation, provides a means for the external regulation of expression of that foreign gene. The method employed for transformation of the plant cells is not especially germane to this invention and any method suitable for the target plant may be employed. Transgenic plants are obtained by regeneration from the transformed cells. Numerous transformation procedures are known from the literature such as agroinfection using Agrobacterium tumefaciens or its Ti plasmid, electroporation, microinjection or plants cells and protoplasts, microprojectile transformation, to mention but a few. Reference may be made to the literature for full details of the known methods.

Neither is the plant species into which the chemically inducible sequence is inserted particularly germane to the invention. Dicotyledonous and monocotyledonous plants can be transformed. This invention may be applied to any plant for which transformation techniques are, or become, available. The present invention can therefore be used to control gene expression in a variety of genetically modified plants, including field crops such as canola, sunflower, tobacco, sugarbeet, and cotton; cereals such as wheat, barley, rice, maize, and sorghum; fruit such as tomatoes, mangoes, peaches, apples, pears, strawberries, bananas and melons; and vegetables such as carrot, lettuce, cabbage and onion. The switch is also suitable for use in a variety of tissues, including roots, leaves, stems and reproductive tissues.

In a particularly preferred embodiment of the present invention, the gene switch of the present invention is used to control expression of genes which confer resistance herbicide resistance and/or insect tolerance to plants.

Recent advances in plant biotechnology have resulted in the generation of transgenic plants resistant to herbicide application, and transgenic plants resistant to insects. Herbicide tolerance has been achieved using a range of different transgenic strategies. One well documented example in the herbicide field is the use the bacterial xenobiotic detoxifying gene phosphinothricin acetyl transferase (PAT) from Streptomyces hydroscopicus. Mutated genes of plant origin, for example the altered target site gene encoding acetolactate synthase (ALS) from Arabidopsis, have been successfully utilised to generate transgenic plants resistant t herbicide application. The PAT and ALS genes have been expressed under the control of strong constitutive promoter. In the field of insecticides, the most common example to-date is the use of the Bt gene.

We propose a system where genes conferring herbicide and/or insect tolerance would be expressed in an inducible manner dependent upon application of a specific activating chemical. This approach has a number of benefits for the farmer, including the following:

1. Inducible control of herbicide and/or insect tolerance would alleviate any risk of yield penalties associated with high levels of constitutive expression of herbicide and/or insect resistance genes. This may be a particular problem as early stages of growth

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where high levels of transgene product may directly interfere with normal development. Alternatively high levels of expression of herbicide and/or insect resistance genes may cause a metabolic drain for plant resources.

- The expression of herbicide resistance genes in an inducible manner allows the
   herbicide in question to be used to control volunteers if the activating chemical is omitted during treatment.
- The use of an inducible promoter to drive herbicide and/or insect resistance genes will 3. reduce the risk of resistance becoming a major problem. If resistance genes were passed onto weed species from related crops, control could still be achieved with the herbicide in the absence of inducing chemical. This would particularly be relevant if 10 the tolerance gene confirmed resistance to a total vegetative control herbicide which would be used (with no inducing chemical) prior to sowing the crop and potentially after the crop has been harvested. For example, it can be envisaged that herbicide resistance cereals, such as wheat, might outcross into the weed wild oats, thus conferring herbicide resistance to this already troublesome weed. A further example is 15 that the inducible expression of herbicide resistance in sugar beet will reduce the risk of wild sugar beet becoming a problem. Similarly, in the field of insect control, insect resistance may well become a problem if the tolerance gene is constitutively expressed. The used of an inducible promoter will allow a greater range of insect resistance control mechanisms to be employed. 20

This strategy of inducible expression of herbicide resistance can be achieved with a pre-spray of chemical activator or in the case of slow acting herbicides, for example N-phosphonomethyl-glycine (commonly known as glyphosate), the chemical inducer can be added as a tank mix simultaneously with the herbicide. Similar strategies can be employed for insect control.

This strategy can be adopted for any resistance confering gene/corresponding herbicide combination, which is, or becomes, available. For example, the gene switch of the present invention can be used with:

- 1. Maize glutathione S-transferase (GST-27) gene (see our International Patent Publication No WO90/08826), which confers resistance to chloroacetanilide herbicides such as acetochlor, metolachlor and alachlor.
- Phosphinotricin acetyl transferase (PAT), which confers resistance to the herbicide commonly known as glufosinate.
- 3. Acetolactate synthase gene mutants from maize (see our International Patent
  Publication No WO90/14000) and other genes, which confer resistance to sulphonyl
  urea and imadazolinones.

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4. Genes which confer resistance to glyphosate. Such genes include the glyphosate oxidoreductase gene (GOX) (see International Patent Publication N. WO92/00377); genes which encode for 5-enolpyruvyl-3-phosphoshikimic acid synthase (EPSPS), including Class I and Class II EPSPS, genes which encode for mutant EPSPS, and genes which encode for EPSPS fusion peptides such as that comprised of a chloroplast transit peptide and EPSPS (see for example EP 218 571, EP 293 358, WO91/04323, WO92/04449 and WO92/06201); and genes which are involved in the expression of CPLyase.

Similarly, the strategy of inducible expression of insect resistance can be adopted for any tolerance confering gene which is, or becomes, available.

The gene switch of the present invention can also be used to controlled expression of foreign proteins in yeast and mammalian cells. Many heterologous proteins for many applications are produced by expression in genetically engineered bacteria, yeast cells and other eucaryotic cells such as mammalian cells.

As well as the obvious advantage in providing control over the expression of foreign genes in such cells, the switch of the present invention provides a further advantage in yeasts and mammalian cells where accumulation of large quantities of an heterologous protein can damage the cells, or where the heterologous protein is damaging such that expression for short periods of time is required in order to maintain the viability of the cells.

Such an inducible system also has applicability in gene therapy allowing the timing of expression of the therapeutic gene to be controlled. The present invention is therefore not only applicable to transformed mammalian cells but also to mammals per se.

A further advantage of the inducible system of the present invention in mammalian cells is that, because it is derived from a insect, there is less chance of it being effected by inducers which effect the natural mammalian steroid receptors.

In another aspect of the present invention the gene switch is used to switch on genes which produce potentially damaging or lethal proteins. Such a system can be employed in the treatment of cancer in which cells are transformed with genes which express proteins which are lethal to the cancer. The timing of the action of such proteins on the cancer cells can be controlled using the switch of the present invention.

The gene switch of the present invention can also be used to switch genes off as well as on. This is useful in disease models. In such a model the cell is allowed to grow before a specific gene(s) is switched off using the present invention. Such a model facilitates the study of the effect of a specific gene(s).

Again the method for producing such transgenic cells is not particularly germane to the present invention and any method suitable for the target cell may be used; such methods are known in the art, including cell specific transformation.

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As previously mentioned, modulation of gene expression in the system appears in response to the binding of the HEcR to a specific control, or regulatory, DNA element. A schematic representation of the HEcR gene switch is shown in Figure 6. For ease of reference, the schematic representation only shows three main domains of the HEcR, namely the transactivation domain, DNA binding domain and the ligand binding domain. Binding of a ligand to the ligand binding domain enables the DNA binding domain to bind to the HRE resulting in expression (or indeed repression) of a target gene.

The gene switch of the present invention can therefore be seen as having two components. The first component comprising the HEcR and a second component comprising an appropriate HRE and the target gene. In practice, the switch may conveniently take the form of one or two sequences of DNA. At least part of the one sequence, or one sequence of the pair, encoding the HEcR protein. Alternatively, the nucleic acid encoding the HEcR can be replaced by the protein/polypeptide itself.

Not only does the switch of the present invention have two components, but also one or more of the domains of the receptor can be varied producing a chimeric gene switch. The switch of the present invention is very flexible and different combinations can be used in order to vary the result/to optimise the system. The only requirement in such chimeric systems is that the DNA binding domain should bind to the hormone response element in order to produce the desired effect.

The glucocorticoid steroid receptor is well characterised and has been found to work well in plants. A further advantage of this receptor is that it functions as a homodimer. This means that there is no need to express a second protein such as the ultraspiracle in order to produce a functional receptor. The problem with the glucocorticoid steroid receptor is that ligands used to activate it are not compatible with agronomic practice.

In a preferred aspect of the present invention the receptor comprises glucocorticoid receptor DNA binding and transactivation domains with a *Heliothis* ligand binding domain according to the present invention. The response unit preferably comprising the glucocorticoid hormone response element and the desired effect gene. In the Examples, for convenience, this effect gene took the form of a reporter gene. However, in non-test or non-screen situations the gene will be the gene which produces the desired effect, for example produces the desired protein. This protein may be a natural or exogenous protein. It will be appreciated that this chimeric switch combines the best features of the glucocorticoid system, whilst overcoming the disadvantage of only being inducible by a steroid.

In another preferred embodiment, the *Heliothis* ligand binding domain is changed, and preferably replaced with a non-*Heliothis* ecdysone receptor ligand binding domain. For example, we have isolated suitable sequences from *Spodoptera exigua*.

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Thus, according to another aspect of the present invention there is provided DNA having the sequence shown in Seq ID No. 6.

According to another aspect of the present invention there is provided DNA having part of the sequence shown in Seq ID No. 6, which encodes for the Spodoptera ecdysone ligand binding domain.

According to another aspect of the present invention there is provided DNA having part of the sequence shown in Seq ID No. 6, which encodes for the Spodoptera ecdysone hinge domain.

The present invention also provides the polypeptides coded for by the above DNA sequences of Seq ID No. 6.

A further advantage with such chimeric systems is that they allow you to choose the promoter which is used to drive the effector gene according to the desired end result. For example, placing the foreign gene under the control of a cell specific promoter can be particularly advantageous in circumstances where you wish to control not only the timing of expression, but also which cells expression occurs in. Such a double control can be particularly important in the areas of gene therapy and the use of cytotoxic proteins.

Changing the promoter also enables gene expression to be up- or down-regulated as desired.

Any convenient promoter can be used in the present invention, and many are known in the art.

Any convenient transactivation domain may also be used. The transactivation domain VP16 is a strong activator from Genentech Inc., and is commonly used when expressing glucocorticoid receptor in plants. Other transactivation domains derived for example from plants or yeast may be employed.

In a preferred embodiment of the present invention, the DNA binding domain is the glucocorticoid DNA binding domain. This domain is commonly a human glucocorticoid receptor DNA binding domain. However, the domain can be obtained from any other convenient source, for example, rats.

According to another aspect of the present invention there is provided a method of selecting compounds capable of being bound to an insect steroid receptor superfamily member comprising screening compounds for binding to a polypeptide or fusion polypeptide of the present invention, and selecting said compounds exhibiting said binding.

According to another aspect of the present invention there is provided a compound selected using the method of the present invention.

According to another aspect of the present invention there is provided an agricultural or pharmaceutical composition comprising the compound of the present invention.

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According to yet another aspect of the present invention there is provided the use of the compound of the present invention as a pesticide, pharmaceutical and/or inducer of the switch. It will be appreciated that such inducers may well be useful as insecticides in themselves.

According to a further aspect of the present invention there is provided a method of producing a protein or peptide or polypeptide comprising introducing into a cell of the present invention, a compound which binds to the ligand binding domain in said cell.

Various preferred features and embodiments of the present invention will now be described by way of non-limiting example with reference to the accompanying examples and figures, in which figures:

Figure 1 (Sequence ID No. 1) shows the DNA sequence amplified from first strand cDNA made from mRNA isolated from *Heliothis virescens* Fourth instar larvae. The underlined sequences refer to the position of the degenerate oligonucleotides. At the 5' end the sequence matches that of the oligonucleotide while at the 3' end 12 nucleotides of the original oligonucleotide are observed;

Figure 2 (Sequence ID No. 2) shows the DNA sequence contained within the clone pSK19R isolated from a random primed cDNA *Heliothis virescens* library; Sequence is flanked by EcoRI sites;

Figure 3 (Sequence ID No. 3) shows the DNA sequence contained within the clone pSK16.1 isolated from a random primed cDNA Heliothis virescens library;

Figure 4 (Sequence ID No. 4) DNA sequence of 5'RACE products (in bold) fused to sequence of clone pSK16.1. The ORF (open reading frame) giving rise to the *Heliothis* virescens ecdysone receptor protein sequence is shown under the corresponding DNA sequence;

Figure 5 (Sequence ID No. 5) shows the protein sequence alignment of the ecdysone receptors DmEcR (*Drosophila melanogaster*), CtEcR (*Chironomus tentans*), BmEcR (*Bombyx mori*), MsEcR (*Manduca sexta*), AaEcR (*Aedes aegipti*) and HvEcR (*Heliothis virescens*). "\*" indicates conserved amino acid residue. "." indicates a conservative amino acid exchange;

Figure 6 shows a model of an embodiment of the glucocorticoid/Heliothis ecdysone chimeric receptor useable as a gene switch;

Figure 7 shows a plasmid map of the clone pcDNA319R. The three other mammalian expression vectors were constructed in the same way and look similar but for the size of the insert:

Figure 8 shows a plasmid map of the reporter construct used to analyse the activity of the *Heliothis virescens* ecdysone receptor;

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Figure 9 is a graph which shows the effect of Muristerone A and RH5992 in reporter activity in HEK293 cells co-transfected with pcDNA3H3KHEcR alone (filled bars) or with  $\alpha$ RXR (stripped bars);

Figure 10 shows a plasmid map of the Maize expression vector containing the Glucocorticoid receptor (HG1 or pMF6HG1PAT);

Figure 11 shows a plasmid map of the maize expression vector containing the chimeric glucocorticoid/Drosophila ecdysone receptor pMF6GREcRS;

Figure 12 shows a plasmid map of the maize expression vector containing the chimeric glucocorticoid/Heliothis ecdysone receptor pMF6GRHEcR;

Figure 13 shows a plasmid map of the plant reporter Plasmid containing the glucocorticoid response elements fused to the -60 S35CaMV promoter fused to GUS, p221.9GRE6;

Figure 14 shows a plasmid map of the plant reporter plasmid containing the glucocorticoid response elements fused to the -46 S35CaMV promoter fused to GUS, p221.10GRE6;

Figure 15 shows a graph showing the effect of Muristerone A and Dexamethasone in Maize AXB protoplasts transformed with pMF6HG1PAT (GR) and p221.9GRE6 (reporter);

Figure 16 shows a graph showing the effect of Muristerone A and Dexamethasone in Maize AXB protoplasts transformed with pMF6GREcRS (effector) and p221.9GRE6 (reporter);

Figure 17 shows a graph showing the effect of Muristerone A and Dexamethasone in Maize AXB protoplasts transformed with pMF6GRHEcR (effector) and p221.9GRE6 (reporter);

Figure 18 shows a graph showing the effect of RH5849 in Maize AXB protoplasts transformed with pMF6GREcRS (effector) and p221.9GRE6 (reporter);

Figure 19 shows a graph showing the effect of RH5992 in Maize AXB protoplasts transformed with pMF6GREcRS (effector) and p221.9GRE6 (reporter);

Figure 20 shows a graph showing the effect of RH5992 in Maize AXB protoplasts transformed with pMF6GRHEcR (effector) and p221.9GRE6 (reporter);

Figure 21 shows a graph which shows the dose response effect of RH5992 in Maize AXB protoplasts transformed with pMF6GRHEcR (effector) and p221.9GRE6 (reporter);

Figure 22 shows a plasmid map of the tobacco expression vector containing the chimeric glucocorticoid/ *Drosophila* ecdysone receptor, pMF7GREcRS;

Figure 23 shows a plasmid map of the tobacco expression vector containing the chimeric glucocorticoid/ Heliothis ecdysone receptor, pMF7GRHEcR;

Figure 24 shows a graph which shows the effect of RH5992 in Tobacco mesophyll protoplasts transformed with pMF6GRHEcR (Effector) and p221.9GRE6 (reporter);

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Figure 25 shows a plasmid map of the mammalian expression vector containing the chimeric glucocorticoid/Heliothis ecdysone receptor, pcDNA3GRHEcR;

Figure 26 shows a plasmid map of the reporter plasmid pSWGRE4;

Figure 27 shows a graph which shows a RH5992 dose response curve of CHO cells transfected with pcDNA3GRHEcR and pSWGRE4;

Figure 28 shows a graph which shows the effect of Muristerone A and RH5992 on HEK293 cells co-transfected with pcDNA3GRHEcR and pSWGRE4;

Figure 29 shows a plasmid map of the binary vector ES1;

Figure 30 shows a plasmid map of the binary vector ES2;

Figure 31 shows a plasmid map of the binary vector ES3;

Figure 32 shows a plasmid map of the binary vector ES4;

Figure 33 shows a plasmid map of the effector construct TEV-B112 made to express the HEcR ligand binding domain in yeast;

Figure 34 shows a plasmid map of the effector construct TEV8 made to express the HEcR ligand binding domain in yeast;

Figure 35 shows a plasmid map of the effector construct TEVVP16-3 made to express the HEcR ligand binding domain in yeast;

Figure 36 shows a plasmid map of the mammalian expression vector containing the chimeric glucocorticoid VP16/Heliothis ecdsysone receptor, pcDNA3GRVP16HEcR;

Figure 37 shows a plasmid map of the maize expression vector containing the chimeric glucocorticoid VP16/Heliothis ecdsysone receptor, pMF6GRVP16HEcR;

Figure 38 shows a plasmid map of the maize expression vector containing the chimeric glucocorticoid VP16/Heliothis ecdsysone receptor, pMF7GRVP16HEcR;

Figure 39 shows a graph which shows the effect of RH5992 in Maize AXB protoplasts transformed with pMF6GRVP16HEcR (effector) and p221.9GRE6 (reporter);

Figure 40 (Sequence ID No. 6) shows the DNA sequence of the hinge and ligand binding domains of the Spodoptera exigua ecdysone receptor;

Figure 41 (Sequence ID No. 7) shows the protein sequence alignment of the *Heliothis* 19R and *Spodoptera* SEcR *Taq* clone hinge and ligand binding domains. "\*" indicates conserved amino acid residue. "." indicates a conservative amino acid exchange;

Figure 42 shows a graph which shows the effect of RH5992 on Tobacco mesophyll protoplasts transformed with pMF7GRHEcR (effector) and either p221.9GRE6 (Horizontal strips) or p221.10GRE6 (vertical strips).

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# Example I - Cloning of the Heliothis Ecdysone Receptor

#### A. Probe generation

The rational behind the generation of the probe to isolate *Heliothis* homologues to the steroid/thyroid receptor superfamily members was based on comparing the sequences of developmentally regulated steroid/thyroid receptor superfamily members. The sequences available showed a highly conserved motif within the DNA binding domain of the RAR and THR (thyroid) receptors. The motifs were used to design degenerate oligonucleotides for PCR amplification of sequences derived from cDNA template produced from tissue expected to express developmentally regulated steroid/thyroid receptor superfamily members (ie. larval tissues).

The sense oligonucleotide is based on the peptide sequence CEGCKGFF which at the DNA level yields an oligonucletide with degeneracy of 32 as shown below:

ZnFA5' 5' TGC GAG GGI TGC AAG GAI TTC TT 3'

The antisense oligonucleotide is based on the reverse complement nucleotide sequence derived from the peptide:

COECRLKK

SR

for which four sets of degenerate oligos were made. Namely:

ZnFA3' 5' TTC TTI AGI CGG CAC TCT TGG CA 3'

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ZnFB3' 5' TTC TTI AAI CGG CAC TCT TGG CA 3'

T ATCA

30 ZnFC3' 5' TTC TTI AGI CTG CAC TCT TGG CA 3'

T ATCA

ZnFD3' 5' TTC TTI AAI CTG CAC TCT TGG CA 3'

T ATCA

The PCR amplification was carried out using a randomly primed cDNA library made from mRNA isolated from 4th and 5th instar *Heliothis virescens* larvae. The amplification

was performed using 10<sup>8</sup> pfus (plaque forming units) in 50mM KCI. 20mM Tris HCl pH 8.4, 15mM MgCl2, 200mM dNTPs (an equimolar mixture of dCTP, dATP, dGTP and dTTP), 100ng of ZnFA5 and ZnF3 mixture. The conditions used in the reaction followed the hot start protocol whereby the reaction mixture was heated to 94°C for 5 minutes after which 1 U of Taq polymerase was added and the reaction allowed to continue for 35 cycles of 93°C for 50 seconds, 40°C for 1 minute and 73°C for 1 minute 30 seconds. The PCR products were fractionated on a 2%(w/v) agarose gel and the fragment migrating between 100 and 200bp markers was isolated and subcloned into the vector pCRII (Invitrogen). The sequence of the insert was determined using Sequenase (USB).

The resulting sequence was translated and a database search carried out. The search recovered sequences matching to the DNA binding domain of the Drosophila ecdysone receptor, retinoic acid receptor and the thyroid receptor. Thus, the sequence of the insert in this plasmid, designated pCRIIZnf, is a Heliothis ecdysone cognate sequence (Figure 1) and was used to screen a cDNA library in other to isolate the complete open reading frame.

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#### B. Library screening

The randomly primed cDNA 4th/5th Instar Heliothis virescens library was plated and replicate filter made from the plates. The number of plaques plated was 500,000. The insert fragment of pCRIIZnf was reamplified and 50ng were end labelled using T4 Polynucleotide Kinase (as described in Sambrook et al 1990).

The filter were prehybridised using 0.25%(w/v) Marvel, 5 X SSPE and 0.1%(w/v) SDS at 42°C for 4 hours. The solution in the filters was ten replaced with fresh solution and the denatured probe added. The hybridisation was carried out overnight at 42°C after which the filter were washed in 6 X SSC + 0.1%(w/v) SDS at 42°C followed by another wash at 55°C. The filter were exposed to X-ray film (Kodak) for 48 hours before processing.

The developed film indicated the presence of one strong positive signal which was plaque purified and further characterised. The lambda ZAP II phage was in vivo excised (see Stratagene Manual) and the sequence determined of the resulting plasmid DNA. The clone known as pSK19R (or 19R) contained a 1.933kb cDNA fragment with an open reading frame of 467 amino acids (Figure 2). pSK19R was deposited with the NCIMB on 20 June 1995 and has been accorded the deposit No NCIMB 40743.

Further analysis of pSK19R revealed that a 340 bp EcoRI fragment mapping at the 5' end of pSK19R has strong and significant similarities to a *Drosophila* cDNA encoding glyceraldehyde-3-phosphate dehydrogenase. In order to isolate the correct 5'end sequence belonging to *Heliothis*, the random primed library was re-screened using a probe containing the 5'end of the pSK19R belonging to *Heliothis* ecdysone receptor. The probe was made by PCR using the sense oligonucleotide HecRH3C (5' aattaagettecaccatgecgttaccaatgecaccacc

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3') and antisense oligonucleotide HecrNdeI (5' cttcaaccgacactcctgac 3'). The PCR was carried out as described by Hirst et al., 1992) where the amount of radioisotope used in the labelling was 50uCi of a <sup>32</sup>P-dCTP and the PCR was cycled for 1 minute at 94°C, 1 minute at 60°C and 1 minute at 72°C for 19 cycles. The resulting 353bp radio labelled DNA fragment was denatured and added to prehybridised filters as described for the isolation of pSK19R. The library filters were made from 15 plates each containing 50000 pfus. The library filters were hybridised at 65°C and washed in 3XSSPE + 0.1%SDS at 65°C twice for 30 minutes each. The filters were further washed with 1XSSPE + 0.1%SDS for 30 minutes and exposed to X-ray film (Kodak) overnight. The film was developed and 16 putative positive plaques were picked. The plaques were re-plated and hybridised under the exact same conditions as the primary screen resulting in only one strong positive. The strong positive was consistently recognised by the probe and was plaque purified and in vivo excised. The resulting plasmid pSK16.1 was sequenced (Seq 1D3) which revealed that the 5' end of the clone extended by 205 bp and at the 3' end by 653 bp and resulting in a DNA insert of 2.5 kb. Conceptual translation of the 205 bp yielded 73 amino acids with high similarity to the Drosophila, Aedes aegipti, Manduca and Bombyx sequences of the ecdsysone receptor B1 isoform. However, the whole of the 5' end sequence is not complete since a Methionine start site was not found with a stop codon in frame 5' of the methionine. In order to isolate the remainder of the 5' end coding sequences a 5'RACE protocol (Rapid Amplification of cDNA Ends) was carried out using the BRL-GIBCO 5'RACE Kit. Two types of cDNA were synthesised where the first one used a specific oligonucleotide: 16PCR2A 5' cagetocaggeegeegateteg3' and the second type used random hexamers (oligonucleotide containing 6 random nucleotides). Each cDNA was PCR amplified using the oligonucleotides anchor primer: BRL-GIBCO 5' cuacuacuacuaggecacgegtegactagtacgggiigggiigggiig 3' and 16PCR2A and cycled for 1 minute at 94°C, 1 minute at 60°C and 1 minute at 72°C for 35 cycles. The reaction conditions were 20mM Tris-HCl (pH8.4), 50mM KCl, 1.5mM MgCl<sub>2</sub>, 400nM of each anchor and 16PCR2A primers, 200mM dNTPs (dATP,dCTP,dGTP and

(Universal Amplification Primer 5' caucaucaucauggecaegegtegaetagtae 3') and 16RACE2:

were made and 1ml was use in a second PCR with oligonucleotides UAP:

(5' aegteaecteagaegagetetecatte 3').

The conditions and cycling were the same as those followed for the first PCR.

Samples of each PCR were run and a Southern blot carried out which was probed with a 5' specific primer:

dTTP) and 0.02 U/ml Taq DNA polymerase. Dilutions of 1:50 of the first PCR reactions

(16PCR1 5' cgctggtataacaacggaccattc 3').

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This primer is specific for the 5' most sequence of pSK16.1 and was hybridised at 55°C using the standard hybridisation buffer. The filter was washed at 55°C 3 times in 3XSSPE + 0.1%SDS and exposed to X-ray film for up to 6 hours. The developed film revealed bands recognised by the oligonucleotide migrating at 100bp and 500bp (relative to the markers). A sample of the PCR reaction (4 in total) was cloned into the pCRII vector in the TA cloning kit (Invitrogen). Analysis of 15 clones from 4 independent PCRs yielded sequence upsteam of pSK16.1 (Figure 4).

Translation of the ORF results in a 575 amino acid protein with high similarity in the DNA and ligand binding domains when compared to the ecdysone receptor sequences of Drosophila, Aedes aegypti, Chironomus tentans, Manduca sexta and Bombyx mori (Figure 5). Interestingly, the N-terminal end of the Heliothis sequence has an in frame methinonine start which is 20 amino acids longer that that reported for Drosophila, Aedes aegypti and Manduca sexta. However, the extended N-terminal end in the Heliothis EcR does not have similarity to that of Bombyx mori. Finally, the C-terminal end of the different B1 isoform ecdysone receptor sequences diverge and do not have significant similarity.

#### C. Northern Blot Analysis

The sequence identified by screening the library is expected to be expressed in tissues undergoing developmental changes, thus mRNA from different developmental stages of H. virescens were was isolated and a northen blot produced. The mRNAs were isolated from eggs, 1st, 2nd, 3rd, 4th and 5th instar larvae, pupae and adults. The northern blot was hybridised with a Ndel/XhoI DNA fragment from pSK19R encompassing the 3'end of the DNA binding domain through to the end of the ligand binding domain. The hybridisation was carried out in 1%(w/v)Marvel, 5X SSPE, 0.1%(w/v) SDS at 65°C for 18 to 24 hours. The filters were washed in 3X SSPE + 0.1%(w/v) SDS and 1X SSPE + 0.1%(w/v) SDS at 25 65°C. The filter was blotted dry and exposed for one to seven days. The gene recognises two transcripts (6.0 and 6.5 kb) which appear to be expressed in all stages examined, however, the levels of expression differ in different stages. It should be noted that the same two transcripts are recognised by probes specific to the DNA binding domain and the ligand binding domain. 30 indicating that the two transcripts arise from the same gene either by alternative splicing or alternative use of polyadenylation sites.

In summary, adult and 5th instar larvae have lower levels of expression while all other tissues have subtantial levels of expression.

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# Example II Expression of Heliothis ecdysone receptor in Mammalian cells

To demonstrate that the cDNA encodes a functional ecdysone receptor, effector constructs were generated containing the HEcR under the control of the CMV (cytomegalovirus) promoter, and the DNA expressed in mammalian cells.

Effector constructs

A first mammalian expression plasmid was constructed by placing a HindIII/NotI pSK19R fragment into the pcDNA3 HindIII/NotI vector resulting in pcDNA319R (Figure 7).

A second effector plasmid was constructed wherein the non-coding region of the cDNA 19R was deleted and a consensus Kozak sequence introduced. The mutagenesis was carried out by PCR amplifying a DNA fragment with the oligo HecRH3C:

5'aattaagcttccaccatgccgttaccaatgccaccgaca 3'
containing a unique HindIII restriction enzyme recognition site followed by the mammalian
Kozak consensus sequence, and HecRNdeI:

5'cttcaaccgacactcctgac 3'.

The resulting 353bp PCR fragment was restriction enzyme digested with HindIII and NdeI, gel purified and ligated with 19R NdeI/NotI fragment into a pcDNA3 HindIII/NotI vector resulting in pcDNA3HecR.

A third effector construct was made with the 5' end sequences of pSK16.1 by PCR. The PCR approach involved PCR amplifying the 5' end sequences using a 5' oligonuclotide containing a HindIII restriction cloning site, the Kozak consensus sequence followed by nucleotide sequence encoding for a Methionine start and two Arginines to be added to the 5' end of the amplified fragment:

(16H3K 5' attaagettgeegeeatgegeegaegetggtataacaaeggaeeatte 3'), the 3' oligonucleotide used was HeerNdel. The resulting fragment was restriction enzyme digested, gel purified and subcloned with an Ndel/Notl 19R fragment into pcDNA3 Ndel/Notl vector. The plasmid was named pcDNA3H3KHEcR.

A fourth effector construct was produced which contains the extended N-terminal end sequence obtained from the 5'RACE experiment. Thus, a PCR approach was followed to introduce the new 5' end sequences in addition to a consensus Kozak sequence and a HindIII unique cloning sequence. The sense oligonucleotide used was RACEH3K:

5' attaagettgeegeeatgteecteggegetegtggatae 3',
while the antisense primer was the same as that used before (HecrNdel). The cloning strategy
was the same as used for the pcDNA3H3KHEcR to give rise to pcDNA3RACEH3KHEcR.

The PCR mutagenesis reactions were carried out in the same manner for all constructs. The PCR conditions used were 1 minute at 94°C, 1 minute at 60°C and 1 minute

at 72°C for 15 cycles. The reactions conditions were 50mM Tris-HCl (pH8.4), 25mM KCl, 200mM dNTPs (dATP, dCTP, dGTP and dTTP), 200nM of each olig nucleotide and 2.5U/Reaction of Taq DNA polymerase. For each construct at least 5 independant PCR reactions were carried out and several clones were sequenced to insure that at least one is mutation free.

#### Reporter construct

The reporter plasmid to be co-transfected with the expression vector contained 4 copies of the Hsp27 ecdysone response element (Riddihough and Pelham, 1987) fused to B-globin promoter and the B-Galactosidase gene. The tandem repeats of the ecdysone response element were synthesised as two complementary oligonucleotides which when annealed produced a double standed DNA molecule flanked by an SpeI site at the 5' end and a ClaI site at the 3' end:

#### Recr3A

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5'ctagtagacaagggttcaatgcacttgtccaataagcttagacaagggttcaatgcacttgtccaatgaattcagacaagggttcaat gcacttgtccaatctgcagagacaagggttcaatgcacttgtccaatat 3'

#### Recr3B

5'cgatattggacaagtgcattgaacccttgtctctgcagattggacaagtgcattgaacccttgtctgaattcattggacaagtgcattgaacccttgtctaagcttattggacaagtgcattgaacccttgtcta 3'.

The resulting 135bp DNA fragment was ligated to the vector pSWBGAL SpeI/ClaI resulting in pSWREcR4 (Figure 8). The co-transfection of the two plasmid should result in B-galactosidase activity in the presence of ligand. The experiment relies upon the presence of RXR (a homologue of ultraspiracle) in mammalian cells for the formation of an active ecdysone receptor.

#### Mammalian transfection methods

Transfections of mammalian cell lines (CHO-K1 Chinese hamster ovary)- ATCC number CCL61 or cos-1 (Monkey cell line) were performed using either calcium phosphate precipitation (Gorman, Chapter 6 of "DNA cloning: a practical approach. Vol 2 D.M. Glover ed/.(1985) IRL Press, Oxford ) or using LipofectAMINE (Gibco BRL Cat. No. 18324-012, following manufacturers instructions). Human Epithelial Kidney 293 cells were transfected using analogous methods.

# Results - Native HECR drives transient reporter gene expression in mammalian cells

Co-transfection of pcDNA3H3KHEcR (Effector) and reporter constructs into Human Epithelial Kidney 293 cells (HEK293) in the presence of either Muristerone A or RH5992 resulted in a 2-3 fold induction of reporter activity compared to the no chemical controls (Figure 9). The HEK293 cells were used since they are known to have constitutive levels of CRXR which have been demonstrated to be necessary for *Drosophila* EcR activation by Muristerone A (Yao., et al., 1993). Moreover, to further investigate the need for RXR

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interactions, a cRXR was co-transfected into HEK293 cells (along with the effector and reporter) resulting in a 9 fold induction of reporter activity compared to the untreated cells (Figure 9). The co-transfection of cRXR with reporter and effector increased by four fold the reporter activity compared to cells transfected with effector and reporter alone. Induction was observed both in the presence of either Muristerone A or RH5992. These data clearly demonstrate that the cDNA HECR encodes a functional ecdysone receptor. Moreover, The ability of HEcR to complex with cRXR and bind Muristerone A or RH5992 provide evidence for the usage of the entire HEcR as a component of a mammalian gene switch. In particular, it offers the advantage of reducing uninduced expression of target gene since ecdysone receptor and response elements are not present in mammalian cells.

# Example III - Chimeric constructs and ligand validation in Maize Protoplasts

In order to apply the ecdysone receptor as an inducible system it was deemed necesary to simplify the requirements of the system by avoiding the need of a heterodimer formation to obtain an active complex. The glucocorticoid receptor is known to form homodimers and chimeric constructs of the glucocorticoid receptor transactivating and DNA binding domains fused to the ecdysone receptor hinge and ligand binding domains have been shown to be active as homodimers in mammalian cells in the presence of Muristerone A (an ecdysone agonist)(Christopherson et al., 1992). However, the chimeric receptor is not responsive to 20-hydroxyecdysone (Christopherson et al., 1992).

The analysis of the activation of the glucocorticoid/Heliothis ecdysone chimeric receptor entailed the production of two other control effector constructs. The first one of the constructs contained the intact glucocorticoid receptor while the second one contained a glucocorticoid/Drosophila ecdysone chimeric receptor.

#### Effector constructs

(i) Glucocorticoid receptor Maize expression construct

The glucocorticoid receptor DNA for the Maize transient expression construct was produced via the polymerase chain reaction (PCR) of Human Fibrosarcoma cDNA (HT1080 cell line, ATCC#CCl121) library (Clontech)(see Hollenberg et al., 1985). The PCR approach taken was to amplify the 2.7kb fragment encoding the glucocorticoid receptor in two segments. The first segment entails the N-terminal end up to and including the DNA binding domain while the second fragment begins with the hinge region (amino acid 500) thought to the end of the reading frame. Thus, the PCR primer for the N-terminal end segment was designed to contain an EcoRI site and the Kozak consensus sequence for translation initiation

GREcoRI 5'attgaattccaccatggactccaaagaatcattaactc 3'.

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The 3'end primer contains a XhoI site in frame with the reading frame at amino acid 500 of the published sequence:

GRXhoI 5' gagactectgtagtggcctcgagcattecttttattttttte 3'.

The second fragment of the glucocorticoid receptor was produced with a 5' end oligonucleotide containing an XhoI site in frame with the open reading frame at the begining of the hinge region (amino acid 500):

GRHinge 5' attetegagatteageaggeeactaeaggag 3'

while the 3' end oligonucleotide contained an EcoRI site 400 bp after the stop codon:

GRStop 5' attgaattcaatgctatcgtaactatacaggg 3'.

The glucocorticoid receptor PCR was carried out using Vent polymerase (Biolabs) under hot start conditions followed by 15 cycles of denaturing (94°C for 1 minute), annealing (66°C for 1 minute) and DNA synthesis (72°C for 3 minute). The template was produced by making first strand cDNA as described in the TA cloning kit (Invitrogen) after which the PCR was carried out in 10mM KCl, 10mM (NH<sub>4</sub>)<sub>2</sub>SO<sub>4</sub>, 20mM TRIS-HCl pH 8.8, 2 mM MgSO<sub>4</sub>, 0.1% (v/v) Triton X-100, 200 mM dNTPs, 100ng of each Primer and 2 U of Vent

Polymerase. The PCR products was restriction enzyme digested with EcoRI and XhoI and subcloned into pBluescript SK (pSK) EcoRI. The resulting plasmid pSKHGI was sequenced and found to lack any mutations from the published sequences (apart from those introduced in the PCR primers) (Hollenberg et al., 1985).

The 2.7kb EcoRI fragment was subcloned into the vector pMF6PAT EcoRI resulting in pMF6HGIPAT (Figure 10).

(ii) Maize expression construct containing a Glucocorticoid/ Drosophila ecdysone chimeric receptor.

The glucocorticoid receptor portion of the chimeric receptor was isolated from pSKHGI by producing a 1.5kb BamHI/XhoI restriction fragment containing the N-terminal end up to and including the DNA binding domain.

The Drosophila ecdysone receptor portion was isolated through PCR of first stand cDNA prepared from Drosophila adult mRNA. The PCR was carried out using a 5' oligonucleotide containing a Sall site (ie. Drosophila ecdysone receptor contains a XhoI site at the end of the ligand binding domain) which starts at the beginning of the hinge region: amino acid 330, Ecr8 attgccacaacggccggaatggccgtcccggag 3'.

The 3' end oligonucleotide contains an BamHI site adjacent to the stop codon: EcRstop 5' tegggettigttaggatectaageegtggtegaatgeteegacttaac 3'.

The PCR was carried out under the conditions described for the amplification of the Glucocorticoid receptor and yielded a 1.6 kb fragment. The fragment was introduced into

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pSK Sall/BamHI and the sequence determined and compared to the published one (Koelle et al., 1991).

The maize transient expression plasmid was produced by introducing into pMF6 BamHI vector the 1.5kb BamHI/XhoI glucocorticoid receptor fragment and the 1.6kb Sall/BamHI *Drosophila* receptor portion to yield the chimeric plasmid pMF6GREcRS (Figure 9).

(iii) Construction of the Glucocorticoid/Heliothis ecdysone chimeric receptor Maize transient expression plasmid.

The Glucocorticoid receptor portion of the chimera was produced as describe in Example II(ii). The production of the *Heliothis* ecdysone receptor portion involves the introduction of a Sall recognition site at the DNA binding/hinge domain junction (amino acid 229). The addition of the Sall site:

Hecrsal 5'attgtcgacaaaggcccgagtgcgtggtgccggag 3'

was achieved via PCR mutagenesis making use of an unique AccI site 107bp downstream of the juction point (or 1007 bp relative to Seq 1D No 4):

Hecracc 5' teacattgeatgatgggaggeatg 3'.

The PCR was carried out using Taq polymerase (2.5 U) in a reaction buffer containing 100ng of template DNA (pSK19R), 100ng of Hecrsal and Hecracc, 20 mM TRIS-HCl pH 8.4, 50mM KCl, 10mM MgCl<sub>2</sub>, 200mM dNTPs. The reaction was carried out with an initial denaturation of 3 minutes followed by 15 cycles of denaturation (1 minute at 94°C), annealing (1 minute at 60°C) and DNA synthesis (1 minute at 72°C). The DNA was restriction enzyme digested and subcloned into pSK Sall/Sacl with the 1.2kb Accl/Sacl 3' end HecR fragment to yield pSK HeCRDEF (or containing the hinge and ligand binding domains of the Heliothis ecdysone receptor). The construction of the maize transient expression plasmid containing the Glucocorticoid/Heliothis ecdysone chimeric receptor involved the ligation of pMF6 EcoRl/Sacl with the 1.5kb EcoRl/Xhol fragment of Glucocorticoid receptor N-terminal end and the 1.2 kb Sall/Sacl fragment of pSk HEcRDEF to yield pMF6GRHEcR (Figure 10). Reporter plasmids

Two reporter plasmids were made by inserting the into p221.9 or p221.10

BamHI/HindIII vectors two pairs or oligonucleotides containing six copies of the glucocorticoid response element (GRE). The two sets of oligonucleotides were designed with restriction enzyme recognition sites so as to ensure insertion of the two pairs in the right orientation. The first oligonucleotide pair GRE1A/B is 82 nucletides long and when annealed result in a DNA fragment flanked with a HindIII site at the 5' end and a SalI site at the 3' end: GRE1A

5'agettegaetgtaeaggatgttetagetaetegagtagetagaacateetgtaeagtegagtagetagaacateetgtaeag 3'

#### **GREIB**

5'tegactgtacaggatgttctagctactegactgtacaggatgttctagctactegagtegctagaacatectgta cagtega 3'.

The second pair of oligonucleotides is flanked by a Sall site at the 5' end and a BamHI site at the the 3' end

GRE2A 5' tegactagetagaacateetgtacagtegagtagetagaacateetgt acagtegagtagetagaacateetgtacag 3'

GRE2B 5'gatectgtacaggatgttetagetactegactgtacaggatgttetagetactegactgtacaggatgttetagetag 3'.

The resulting plasmids were named p221.9GRE6 (Figure 13) and p221.10GRE6 (Figure 14)(used in later Example). The difference between p221.9 and p221.10 plasmids is that p221.9 contains the -60 35SCaMV minimal promotor while p221.10 (p221.10GRE6) contains the -46 35SCaMV minimal promotor.

#### Method

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Protoplasts were isolated from a maize suspension culture derived from BE70 x A188 embryogenic callus material, which was maintained by subculturing twice weekly in MS0.5<sub>mod</sub> (MS medium supplemented with 3% sucrose, 690mg/l proline, 1g/l myo-inositol, 0.2g/l casein acid hydrolysate, 0.5mg/1 2,4-D, pH5.6). Cells from suspensions two days post subculture were digested in enzyme mixture (2.0% Cellulase RS, 0.2% Pectolyase Y23, 0.5M Mannitol. 5mM CaCl<sub>2</sub>2H<sub>2</sub>O, 0.5% MES, pH5.6, ~660mmol/kg) using ~10ml/g cells, incubating at 25°C, dim light, rotating gently for -2 hours. The digestion mixture was sieved sequentially through 250 um and 38 µm sieves, and the filtrate centrifuged at 700 rpm for 3.5 minutes, discarding the supernatant. The protoplasts were resuspended in wash buffer (0.358M KCl, 1.0mM NHLNO<sub>3</sub>, 5.0mM CaCl<sub>2</sub>2H<sub>2</sub>O, 0.5mM KH<sub>2</sub>PO<sub>4</sub>, pH4.8, ~670mmol/kg) and pelleted as before. This washing step was repeated. The pellet was resuspended in wash buffer and the protoplasts were counted. Transformation was achieved using a Polyethylene glycol method based on Negrutiu et.al. Protoplasts were resuspended at 2 x 10<sup>6</sup>/ml in MaMg medium (0.4M Mannitol. 15mM MgCl2. 0.1% MES, pH5.6, -450mmol/kg) aliquotting 0.5ml / treatment (i.e. 1x10<sup>6</sup> protoplests / treatment). Samples were heat shocked at 45°C for 5 minutes then cooled to room temperature. 10ug each of p221.9GRE6 and pMF6HR1PAT (GR) (1mg/ml)/ treatment were added and mixed in gently, followed by immediate addition of 0.5ml warm (~45°C) PEG solution (40% PEG 3,350MW in 0.4M Mannitol, 0.1M Ca(NO<sub>3</sub>)<sub>2</sub>, pH8.0), which was mixed in thoroughly but gently. Treatments were incubated at room temperature for 20-25 minutes, then 5ml 0.292M KCl (pH5.6, ~530mmol/kg) was added step-wise, 1ml at a time, with mixing. The treatments were incubated for a further 10-15 minutes prior to pelleting the protoplasts by centrifuging as before. Each protoplast treatment was resuspended in 1.5ml culture medium (MS medium, 2% sucrose, 2mg/l 2,4-D, 9% Mannitol, pH5.6, ~700mmol/kg) +/- 0.0001M dexamethasone (glucocorticoid). The samples were incubated in 3cm dishes at 25°C, dark, for 24-48 hours prior to harvesting. Fluorometric

assays for GUS activity were performed with the substrate 4-methylumbelliferyl-D-glucuronide using a Perkin-Elmer LS-35 fluorometer (Jefferson et al., 1987). Protein concentration of tissue homogenates were determined by the Bio-Rad protein assay (Bradford, 1976). The method was repeated for each effector construct.

#### 5 Results

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## Reporter gene assay

A reporter gene construct (p221.9GRE6) was generated containing the GUS reporter gene under the control of a -60 CaMV 35S promoter with 6 copies of the glucocorticoid response element. To test this construct was functional in maize protoplasts a cotransformation assay was performed with the reporter construct p221.9GRE6 and the effector construct pMF6HR1PAT (GR) construct containing the entire glucorticoid receptor.

Figure 15 shows that Reporter p221.9GRE6 alone or reporter plus effector pMF6HR1PAT (GR) with no activating chemical gave no significant expression. When reporter plus effector were co-transformed into maize protoplasts in the presence of 0.0001M dexamethasone (glucocorticoid), a significant elevation of marker gene activity was observed (Figure 15). The response is specific to glucorticoid as the steroid Muristerone A does not lead to induced levels of expression. These studies clearly show the reporter gene construct p221.9GRE6 is capable of monitoring effector /ligand mediated gene expression. Chimeric ecdysone effector constructs mediate inducible expression in maize transient protoplasts assays

A chimeric effector plasmid pMF6GREcRS was constructed, containing the ligand binding domain from the *Drosophila* ecdysone receptor and the DNA binding and transactivation domain from the glucorticoid receptor. To confirm the reporter gene construct p221.9GRE6 could respond to a chimeric ecdysone effector construct, a series of co-transformation into maize protoplasts was performed.

Figure 16 shows that reporter (p221.9GRE6) alone or reporter plus effector (pMF6GREcRS) with no activating chemical, gave no significant expression in maize protoplasts. When reporter plus effector were co-transformed into maize protoplasts in the presence of 100µM Muristerone A, a significant elevation of marker gene activity was observed. The response was specific to Muristerone A, as the steroid dexamethasone did not lead to induced levels of expression. These studies clearly showed the reporter gene construct p221.9GRE6 is capable of monitoring chimeric ecdysone effector /ligand mediated gene expression.

A second chimeric effector construct pMF6GRHEcR, was generated containing the ligand binding domain from *Heliothis* ecdysone receptor. When co-transformed into maize protoplasts with the reporter plasmid p221.9GRE6, no response to 100µM Muristerone or

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100µM dexamethasone was observed (Figure 17). These data clearly show the *Drosophila* and *Heliothis* ligand binding domains exhibit different properties.

When the effector plasmid pMF6GREcRS, containing the ligand binding domain from *Drosophila*, was tested with the reporter p221.9GRE6 in presence of the non-steroidal ecdysone agonists RH5849 and RH5992 (mimic), no chemical induced reporter gene activity was observed (Figures 18 and 19).

When the effector plasmid pMF6GRHEcR, containing the ligand binding domain from *Heliothis*, was tested with the reporter p221.9GRE6 in presence of the non-steroidal ecdysone agonists RH5992 (mimic), significant chemical induced reporter gene activity was observed (Figure 20). These data demonstrate the ligand binding domain from *Heliothis* has different properties to the *Drosophila* receptor in that the former responded to the non-steroidal ecdysteroid agonist RH5992. Figure 21 demonstrates the effector plasmid pMF6GRHEcR confers RH5992 dependant inducibility on the reporter p221.9GRE6 in a dose responsive manner. Induction was observed in a range from 1µM-100µM RH5992.

## Example IV - Testing of effector vectors in Tobacco protoplasts

The experiments carried out in the previous example demonstrated the specific effect of RH5992 (mimic) on pMF6GRHEcR in maize protoplasts. It is the aim in this example to show the generic application to plants of the glucocorticoid/Heliothis ecdysone chimeric receptor switch system. Tobacco shoot cultures cv. Samsun, were maintained on solidified MS medium + 3% sucrose in a controlled environment room (16 hour day / 8 hour night at 25°C, 55% R.H), were used as the source material for protoplasts. Leaves were sliced parallel to the mid-rib, discarding any large veins and the slices were placed in CPW13M 13% Mannitol, nH5.6, ~860mmol/kg) for ~1 hour to pre-plasmolyse the cells. This solution was replaced with enzyme mixture (0.2% Cellulase R10, 0.05% Macerozyme R10 in CPW9M (CPW13M but 9% Mannitol), pH5.6, ~600mmol/kg) and incubated in the dark at 25°C overnight (~16 hours). Following digestion, the tissue was teased apart with forceps and any large undigested pieces were discarded. The enzyme mixture was passed through a 75µm sieve and the filtrate was centrifuged at 600rpm for 3.5 minutes, discarding the supernatant. The pellet was resuspended in 0.6M sucrose solution and centrifuged at 600rpm for 10 minutes. The floating layer of protoplasts was removed using a pasteur pipette and diluted with CPW9M (pH5.6, ~560mmol/kg). The protoplasts were again pelleted by centrifuging at 600rpm for 3.5 minutes, resuspended in CPW9M and counted. A modified version of the PEG-mediated transformation above was carried out. Protoplasts were resuspended at 2x10<sup>6</sup>/ml in MaMg medium and aliquotted using 200µl / treatment (i.e. 4x10<sup>5</sup> protoplasts / treatment). 20µg each of pMF6GRHEcRS and p221.9GRE6 DNA (1mg/ml) were added

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followed by 200µl PEG solution and the solutions gently mixed. The protoplasts were left to incubate at room temperature for 10 minutes before addition of 5ml MSP19M medium (MS medium, 3% sucrose, 9% Mannitol, 2mg/l NAA, 0.5mg/l BAP, pH5.6, ~700mmol/kg) +/- 10 µM RH5992. Following gentle mixing, the protoplasts were cultured in their tubes, lying horizontally at 25°C, light. The protoplasts were harvested for the GUS assay after ~24 hours. Effector construct

(i) Construction of a Dicotyledonous expression vector

The vector produced is a derivative of pMF6. pMF6GREcRS was restriction enzyme digested with PstI to produce 3 fragments namely, 3.4(Adh Intronless pMF6), 3.2(GREcRS) and 0.5(Adh intron I) kb). Isolation and religation of the 3.4 and 3.2 kb fragments resulted in pMF7GREcRS (Figure 22). pMF7GREcRS was restriction enzyme digested with EcoRI/SacI resulting in the 3.4kb pMF7 EcoRI/SacI vector which when isolated and purified was ligated to a 1.5 kb EcoRI/XhoI N-terminal end of the glucocorticoid receptor and the 1.2 kb Sall/SacI Heliothis ecdysone C-terminal end sequences to produce pMF7GRHEcR (Figure 23).

#### Reporter plasmid

The reporter plasmids constructed for the maize transient experiments were the same as those used without alteration in the tobacco leaf protoplast transient expression experiments.

20 Results - Chimeric ecdysone effector constructs mediate inducible expression in tobacco transient protoplast assays

Experiments were performed to demonstrate that the effector plasmid pMF6GRHEcR can confer chemical dependant inducible expression on the reporter p221.9GRE6 in tobacco mesophyll protoplasts.

Figure 24 shows that reporter (p221.9GRE6) alone or reporter plus effector (pMF7GRHEcR) with no activating chemical, gave no significant expression in tobacco protoplasts. When reporter plus effector were co-transformed into tobacco protoplasts in the presence of 10µM RH5992, a significant elevation of marker gene activity was observed. These data show a chimeric ecdysone effector construct, containing the *Heliothis* ligand binding domain can confer non-steroidal ecdysteroid dependant expression on reporter gene constructs in both monocotyledonous and dicotyledonous species.

#### Example V - Chimeric activity in Mammalian cells

#### Effector constructs

5 (i) Construction of Glucocorticoid/Heliothis ecdysone chimeric receptor.

The mammalian expression vector used in this experiment was pcDNA3 (Invitrogen). The GRHEcR 2.7kb BamHI DNA fragment (isolated from pMF6GRHEcR) was introduced into the pcDNA3 BamHI vector. The recombinants were oriented by restriction enzyme mapping. The DNA sequence of the junctions was determined to ensure correct orientation and insertion (pcDNA3GRHEcR, Figure 25).

#### Reporter construct

The reporter plasmid for mammalian cell system was produced by taking pSWBGAL plasmid and replacing the CRESW SpeI/ClaI fragment for a synthetic 105 bp DNA fragment containing 4 copies of the glucocorticoid response element (GRE) and flanked by SpeI at the 5' end and Af1II at the 3' end.

The oligonucleotides were synthesised using the sequences:

#### **GREspel**

5'ctagttgtacaggatgttctagctactcgagtagctagaacatcctgtacagtcgagtagctagaacatcctgtacagtcgagtagct agaacatcctgtacac 3'

#### 20 GREaf12

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5'ttaagtgtacaggatgttctagctactcgactgtacaggatgttctagctactcgactgtacaggatgttctagctactcgagtagctaggatagttctagctactcgagtagctaggatgttctagctactcgagtagctaggatgttctagctactcgagtagctaggatgttctagctactcgagtagctaggatgttctagctactcgagtagcta

The two oligonucleotides were purified annealed and ligated to pSWBGAL SpeI/AfIII to produce pSWGRE4 (Figure 26).

25 Results - Chimeric HEcR drives transient reporter gene expression in mammalian cells

No expression was detected when a reporter gene construct pSWGRE4, comprising of a minimal β-globin promoter containing four copies of the glucocorticoid response element, fused to a β-galactosidase reporter gene, was introduced into CHO cells. Similarly, no expression was detected when pSWGRE4 and an effector plasmid pCDNA3GRHEcR, containing the transactivation and DNA binding domain from the glucocorticoid receptor and the ligand binding domain from the *Heliothis* ecdysone receptor, under the control of the CMV promoter were co-transformed into CHO-K1 or HEK293 cells. When co-transformed CHO (Figure 27) and HEK293 cells (Figure 28) were incubated in the presence of the non-steroidal ecdysone agonists RH5992 (mimic), significant chemical induced reporter gene activity was observed. Equally, induction of reporter activity was observed when HEK293 cells transfected with pcDNA3GRHEcR and reporter were treated with Muristerone A (Figure 28).

# Example VI - Screening system allows new chemical activators and modified ligand binding domains to be tested in Mammalian cells

The basis of a screening system are in place after the demonstration that the chimeric receptor was activated in the presence of RH5992. A screen was carried out using CHO cells transiently transfected with both pSWGRE4 (reporter) and pcDNA3GRHEcR (effector) constructs. In the first instance 20 derivatives compounds of RH5992 were screened. It was observed that 7 out of the 20 compounds gave an increased reporter gene activity compared to untreated cells. A second screen was carried out in which 1000 randomly selected compounds were applied to transiently transfected CHO cells. Two compounds were found to activate reporter gene activity above that from the untreated controls. The second screen suggest that this cell based assay is a robust and rapid way to screen a small library of compounds, where a thousand compounds can be put through per week.

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#### Example V - Stably transformed Tobacco plants

#### Stable Tobacco vectors

The components of the stable Tobacco vectors were put together in pBluescript prior to transfer into the binary vector. The production of stable transformed plants entails the production of a vector in which both components of the switch system (ie. effector and reporter) are placed in the same construct to then introduce into plants.

The methodology described below was used to produce four different stable Tobacco vectors. The method involves three steps:

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 pBluescript SK HindIII/EcoRI vector was ligated to either GRE6-4635SCaMVGUSNOS HindIII/EcoRI (from p221.10GRE6) or GRE6-6035SCaMVGUSNOS HindIII/EcoRI (from p221.9GRE6) resulting in plasmid pSK-46 and pSK-60.

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2. This step involves the addition of the chimeric receptor (35SGRHEcRNOS or 35SGRVP16HEcRNOS) to pSK-60 or pSK-46. Thus a pSK-60 (or pSK-46) XbaI vector was ligated with either the 3.4kb 35SGRHEcRNOS XbaI or the 3.0kb 35SGRVP16HEcRNOS XbaI DNA fragment to produce pSKES1 (pSKGRE6-6035SCaMVGUSNOS-35SGRHEcRNOS), pSKES2 (pSKGRE6-4635SCaMVGUSNOS-35SGRHEcRNOS), pSKES3 (pSKGRE6-6035SCaMVGUSNOS-35SGRVP16HEcRNOS) and pSKES4 (pSKGRE6-4635SCaMVGUSNOS-35SGRVP16HEcRNOS).

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- 3. Transfer from pBluescript based vectors to binary vectors. The transfer of the ES1 (Figure 29) ES2 (Figure 30), ES3 (Figure 31) or ES4 (Figure 32) DNA fragments into the binary vector JR1 involves five steps:
- (i) Restriction enzyme digestion of pSKES1 (ES2, ES3, and ES4) with ApaI and NotI to liberate the insert from the vector pBluescript.
- (ii) The two DNA fragments were BamHI methylated for 2 hours at 37°C in TRIS-HCl, MgCl, 80uM SAM (S-adenosylmethionine) and 20 U of BamHI methylase.
- 10 (iii) Ligate a Apal/NotI linker onto the fragment. The linker was designed to have an internal BamHI site:

ApaBNot1 5' cattggatecttage 3' and

ApaBNot2 5'ggccgctaaggatccaatgggcc 3'.

- (iv) Restriction enzyme digest the protected and linkered fragment with BamHI and fractionate the products on a 1%(w/v) Agarose gel. The protected DNA fragment (5.5kb) was cut out of the gel and purified.
  - (v) A ligation of JRI BamHI vector with the protected band was carried out to produce JRIESI (JRIES2, JRIES3 or JRIES4). The DNA of the recombinant was characterised by restriction mapping and the sequence of the junctions determined.

The plant transformation construct pES1, containing a chimeric ecdysone receptor and a reporter gene cassette, was transferred into Agrobacterium tumefaciens LBA4404 using the freeze/thaw method described by Holsters et al. (1978). Tobacco (Nicotiana tabacum cv Samsun) transformants were produced by the leaf disc method (Bevan, 1984). Shoots were regenerated on medium containing 100mg/l kanamycin. After rooting, plantlets were transferred to the glasshouse and grown under 16 hour light/8 hour dark conditions.

Results - Chimeric ecdysone effector constructs mediate inducible expression in stably tobacco plants

Transgenic tobacco plants were treated in cell culture by adding 100µM RH5992 to MS media. In addition seedlings were grown hydroponically in the presence or absence of RH5992. In further experiments 5mM RH5992 was applied in a foliar application to 8 week old glasshouse grown tobacco plants. In the three methods described uninduced levels of GUS activity were comparable to a wild type control, while RH5992 levels were significantly elevated.

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# **Ecdysone switch modulation and optimisation**

# Example VIII - Yeast indicator strains for primary screen of chemical libraries

A set of yeast indicator strains was produced to use as a primary screen to find chemicals which may be used in the gene switch. The properties of the desired chemicals should include high affinity resulting in high activation but with different physico-chemical characteristics so as to increase the scope of application of the technology. Moreover, the production of this strain also demonstrates the generic features of this switch system. Effector vector

A base vector for yeast YCp15Gal-TEV-112 was generated containing:

Backbone - a modified version of pRS315 (Sikorski and Hieter (1989) Genetics 122, 19-27)
a shuttle vector with the LEU2 selectable marker for use in yeast;

- 15 ADH1 promoter (BamHI- Hind III fragment) and ADH1 terminator (Not I- Bam HI fragment) from pADNS (Colicelli et al PNAS 86, 3599-3603);

  DNA binding domain of GAL4 (amino acids 1-147; GAL4 sequence is Laughon and Gesteland 91984) Mol. Cell Biol. 4, 260-267) from pSG424 (Sadowski and Ptashne (1989) Nuc. Acids Res. 17, 7539);
- Activation domain an acidic activation region corresponding to amino acids 1-107 of activation region B112 obtained from plasmid pB112 (Ruden et al (1991) Nature 350, 250-252).

The plasmid contains unique Eco RI, Nco I and Xba I sites between the DNA binding domain and activation domains.

Into this vector a PCR DNA fragment of the *Heliothis* ecdysone receptor containing the hinge, ligand binding domains and the C-terminal end was inserted. The 5' oligonneleotide is flanked by an Neol restriction recognition site and begins at amino acid 259: HecrNeol 5' aattecatggtacgacgacagtagacgatcac 3'.

The 3' oligonucleotide is flanked by an Xbal site and encodes for up to amino acid

HecRXbaI 5' ctgaggtctagagacggtggcggcggcc 3'.

The PCR was carried out using vent polymerase with the conditions described in Example IA. The fragment was restriction enzyme digested with NcoI and XbaI purified and ligated into YCp15GALTEV112 NcoI/XbaI vector to produce YGALHeCRB112 or TEV-B112 (Figure 34). In order to reduce constitutive activity of the YGALHeCRB112 plasmid a YGALHeCR plasmid was produced in which the B112 activator was deleted by restriction enzyme digesting YGALHeCRB112 with XbaI/SpeI followed by ligation of the resulting

vector (ie. SpeI and XbaI sites when digested produce compatible ends)(TEV-8, Figure 33). An effector plasmid was constructed whereby the B112 transactivating domain was excised from YGalHecRB112 with XbaI and replaced with the VP16 transactivation domain DNA fragment (encoding amino acids 411 and 490 including the stop codon). The resulting vector was named YGalHecRVP16 or TEVVP16-3 (Figure 35).

#### Reporter construction for yeast

The S. cerevisiae strain GGY1::171 (Gill and Ptashne (1987) Cell 51, 121-126), YT6::171 (Himmelfarb et al (1990) Cell 63, 1299-1309) both contain reporter plasmids consisting of the GALA-responsive GAL1 promoter driving the E. coli B-galactosidase gene. These plasmids are integrated at the URA3 locus. The reporter strain YT6::185 contains the reporter plasmid pJP185 (two synthetic GALA sites driving the B-galactosidase gene) integrated at the URA3 locus of YT6 (Himmelfarb et al). (Note-the parental strains YT6 and GGY1 have mutations in the GALA and GAL80 genes, so the reporter genes are inactive in the absence of any plasmids expressing GALA fusions).

#### 15 Yeast assay

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Standard transformation protocols (Lithium acetate procedure) and selection of colonies by growth of cells on selective media (leucine minus medium in the case of the YCp15Gal-TEV-112 plasmid)- as described in Guthrie and Fink )1991) Guide to Yeast Genetics and Molecular Biology: Methods in Enzymology Vol. 194 Academic Press) and the reporter gene assay is a modification of that described in Ausabel et al (1993) Current Protocols in Molecular Biology (Wiley) Chapter 13).

Results - Automated screening system allows new chemical activators and modified ligand binding domains to be tested in yeast

An effector vector pYGALHEcRB112 has been generated containing a GAL4 DNA binding domain, a B112 activation domain and the ligand binding region from *Heliothis virescens*. In combination with a GAL reporter vector, pYGALHEcRB112 form the basis of a rapid, high throughput assay which is cheap to run. This cell-based assay in yeast (*Saccharomyces cerevisiae*) will be used to screen for novel non-steroidal ecdysone agonists which may of commercial interest as novel insecticides or potent activators of the ecdysone gene switch system. The demonstration of an efficient system to control gene expression in a chemical dependant manner, forms the basis of an inducible system for peptide production in yeast.

The yeast screening system forms the basis of a screen for enhanced ligand binding using the lac Z reporter gene vector to quantitatively assay the contribution of mutation in the ligand binding domain. Alternatively, enhanced ligand binding capabilities or with a selection cassette where the lac Z reporter is replaced with a selectable marker such as uracil (URA 3), tryptophan (Trp1) or leucine (Leu2), and histidine (His). Constructs based on

pYGALHECRB112 with alterations in the ligand binding domain are grown under selection conditions which impair growth of yeast containing the wild type ligand binding domain. Those surviving in the presence of inducer are retested and then sequenced to identify the mutation conferring resistance.

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#### Example IX - Optimisation of chimeric receptor using a strong transactivator

Construction of mammalian expression plasmid with chimeric receptor containing Hernex Simplex VP16 protein sequences.

The construction of this chimeric receptor is based on replacing the sequences encoding for the glucocorticoid receptor transactivating domain with those belonging to the VP16 protein of Herpex simplex. Thus PCR was used to generate three fragments all to be assembled to produce the chimeric receptor. The PCRs were carried out as described in Example II, iii. The first fragment includes the Kozak sequences and methionine start site of the glucocorticoid receptor to amino acid 152 of the glucocorticoid receptor.' The oligonucleotides used for the generation of this fragment included an EcoRI site at the 5' end: GR1A 5' atatgaattccaccatggactccaaagaatc 3' and at the 3' end a NheI restriction enzyme recognition site:

GR1B 5' atat<u>ectase</u>tgtgggggcagcagagagagagagggg 3'.

The second fragment also belongs to the glucocorticoid receptor and begins with a Nhel site in frame with amino acid 406:

GR2A 5'atatgctagctccagctcctcaacagcaacaac 3'

and ends with a XhoI site at amino acid 500:

GR2B 5'atatctcgagcaattccttttatttttttc 3'.

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The two fragments were introduced into pSKEcoRI/SacI in a ligation containing GR1A/B EcoRI/NheI, GR2A/B NheI/XhoI and HEcR SaII/SacI (from pSKHEcRDEF) to yield pSKGRDHEcR. The GR sequences and junctions of the ligation were found to be mutation free.

The third fragment to be amplified was a sequence between amino acid 411 to 490 of the herpes simplex VP16 protein. The amplified fragment was flanked with SpeI recognition sites. Spel produces compatible ends to those of Nhel sites. The oligonucleotides used:

VP16C 5' attactagttctgcggecccccgaccgat 3' and

VP16E 5' aattactagtcccaccgtactcgtcaattcc 3'

produced a 180 bp fragment which was restriction enzyme digested with SpeI and introduced into pSKGRAHECR NheI vector to produce pSKGRVP16HECR. The DNA from the latter was sequenced and and f und to be mutation free, the junctions were also shown to be in frame with those of the glucocorticoid receptor.

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The 2.2 kb EcoRV/NotI GRVP16HEcR fragment was introduced into a pcDNA3 EcoRV/NotI vector resulting in pcDNA3GRVP16HEcR (Figure 36).

Construction of plant transient expression effector plasmids containing the chimeric receptor with VP16 sequences

The same procedure was carried out to clone the GRVP16HeCR DNA fragment into tobacco(pMF7b) and maize(pMF6) expression vectors. A 2.2kb BamHI DNA fragment was isolated from pcDNA3GRVP16HeCR and ligated in to the pMF6 BamHI (or pMF7b BamHI) vector to produce pMF6GRVP16HeCR (Figure 37) (or pMF7GRVP16HeCR) (Figure 38). Results - Addition of strong activation domains enhance ecclysone switch system

The VP16 transactivation domain from herpes simplex virus has been added to a maize protoplast vector pMF6GRHEcR to generate the vector pMF6GRVP16HEcR. When co-transformed into maize protoplasts with the reporter construct p221.9GRE6, in the presence of 100µM RH5992, enhanced levels of expression were seen over pMF6GRHEcR. Figure 39, shows that RH5992 is able to induce GUS levels comparable to those observed with the positive control (p35SCaMVGUS), moreover, a dose response effect is observable.

VP16 enhanced vectors (pES3 and pES4) have been generated for stable transformation of tobacco. Following transformation transgenic progeny containing pES3 and pES4, gave elevated GUS levels following treatment with RH5992, relative to comparable transgenic plants containing the non-VP16 enhanced vectors pES1 and pES2.

An enhanced mammalian vector pcDNA3GRVP16HEcR was prepared for transient transfection of mammalian cell lines. Elevated reporter gene activities were obtained relative to the effector construct (pCDNA3GRHEcR) without the VP16 addition.

"Acidic" activation domains are apparently "universal" activators in eukaryotes (Ptashne (1988) Nature 335 683-689). Other suitable acidic activation domains which have been used in fusions are the activator regions of GAL4 itself (region I and region II; Ma and Ptashne (Cell (1987) 48, 847-853), the yeast activator GCN4 (Hope and Struhl (1986) Cell 46, 885-894) and the herpes simplex virus VP16 protein (Triezenberg et al (1988) Genes Dev. 2, 718-729 and 730-742).

Other acidic and non-acidic transcriptional enhancer sequences for example from plant fungal and mammalian species can be added to the chimeric ecdysone receptor to enhance induced levels of gene expression.

Chimeric or synthetic activation domains can be generated to enhance induced levels of gene expression.

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## Example X - Optimisation by replacement of *Heliothis* ligand binding domain in chimeric effector for that of an ecdysone ligand binding domain of another species

Mutagenesis of the ecdysone ligand binding domain results in the increased sensitivity of the chimeric receptor for activating chemical. This can be achieved by deletions in the ligand binding domain, use of error prone PCR (Caldwell et al., PCR Meth. Applic 2, 28-33 1992), and in vitro DNA shuffling PCR (Stemmer, Nature 370, 389-391 1994). To enhance the efficacy of the listed techniques we have developed a screening system for enhanced levels of induced expression (see below).

An alternative strategy to the mutation of a known ligand binding domain is to identify insect species which are particularly sensitive to ecdysteroid agonists. For example Spodoptera exigua is particularly sensitive to RH 5992. To investigate the role of the ecdysone receptor ligand binding domain in increased sensitivity to RH5992 we have isolated corresponding DNA sequences from of S. exigua (Figure 40, Sequence ID No. 6). Figure 41, Sequence ID No. 7 shows a protein alignment of the hinge and ligand binding domains of the Heliothis virescens and Spodoptera exigua ecdysone receptors. The protein sequence between the two species is well conserved.

Results - Manipulation of the ligand binding domain leads to enhanced induced expression

Isolation of an ecdysone ligand binding domain from another lepidopteran species was carried out by using degenerate oligonucleotides and PCR of first strand cDNA (Perkin Elmer, cDNA synthesis Kit) of the chosen species. The degenerate oligonucleotides at the 5' end were HingxhoA and B and at the 3' end ligandxA/B

- 25 HingxhoA 5' attgctcgagaaagiccigagtgcgtigticc 3'
  - a t
  - HingxhoB 5' attgctcgagaacgiccigagtgtgtigticc 3'
    - a C
- 30 LigandxA 5' ttactcgagiacgtcccaiatctcttciaggaa 3'
  - a tc a
  - ligandxB 5' ttactcgagiacgtcccaiatctcctciaagaa 3'

tta

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RNA was extracted from 4th instar larvae of Spodoptera exigua since Spodoptera exigua appears to be more sensitive to RH5992 than Heliothis (Smagghe and Degheele,

1994). The first strand cDNA was used in PCR reactions under the following conditions 20mM Tris-HCL (pH8.4), 50mM KCl, 1.5mM MgCl<sub>2</sub>, 200mM dNTPs (dATP,dCTP,dGTP and dTTP) and 0.02 U/ml Taq DNA polymerase and in the presence of 1ug of each Hinge (5' 3') and Ligand (5'3') oligonucleotides. The PCR cycling conditions were 94°C for 1 minute, 60°C for 2 minutes and 72°C for 1 minute and 35 cycles were carried out. A sample of the completed reaction was fractionated in a 1% agarose (w/v) 1 x TBE gel, and the resulting 900 bp fragment was subcloned into pCRII vector (Invitrogen). The resulting clone (pSKSEcR 1-10) were further characterised and sequenced.

### 10 Example X - Manipulation of reporter gene promoter regions can modulate chemical induced expression

The context of the effector response element in the reporter gene promoter can be used to modulate the basal and induced levels of gene expression. Six copies of the glucorticoid response element were fused to 46 bp or 60 bp of the CaMV 35S promoter sequence. When used with the effector construct pMF7GRHEcRS the reporter gene construct containing 46 bp of the CaMV 35S promoter gave reduced basal and induced levels of GUS expression relative to the 60 bp reporter construct (Figure 42).

Constructs for plant transformation (pES1 and ES2) have been generated to demonstrate the size of minimal promoter can be used to modulate the basal and induced levels of gene expression in plants.

The number and spacing of response elements in the reporter gene promoter can be adjusted to enhance induced levels of trans-gene expression.

The utility of a two component system (effector and reporter gene cassettes) allows the spatial control of induced expression. Trans-gene expression can be regulated in an tissue specific, organ specific or developmentally controlled manner. This can be achieved by driving the effector construct from a spatially or temporally regulated promoter.

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#### SEQUENCE LISTING

_	(1) GENERAL INFORMATION:	
5	(i) APPLICANT: (A) NAMB: ZENECA LIMITED	
	(B) STREET: 15 STANHOPE GATE (C) CITY: LONDON	
10	(E) COUNTRY: UK (F) POSTAL CODE (ZIP): WIY 6LN	
	(ii) TITLE OF INVENTION: A GENE SWITCH	
15	(iii) NUMBER OF SEQUENCES: 7	
20	<pre>(iv) COMPUTER READABLE FORM:     (A) MEDIUM TYPE: Floppy disk     (B) COMPUTER: IBM PC compatible     (C) OPERATING SYSTEM: PC-DOS/MS-DOS     (D) SOFTWARE: PatentIn Release #1.0, Version #1.30 (EPO)</pre>	
25	(vi) PRIOR APPLICATION DATA:  (A) APPLICATION NUMBER: GB 9510759.5  (B) FILING DATE: 26-MAY-1995	
30	(vi) PRIOR APPLICATION DATA:  (A) APPLICATION NUMBER: GB 9513882.3  (B) FILING DATE: 07-JUL-1995	
<b></b>	<ul><li>(vi) PRIOR APPLICATION DATA:</li><li>(A) APPLICATION NUMBER: GB 9517316.7</li><li>(B) FILING DATE: 24-AUG-1995</li></ul>	
35 ·	(vi) PRIOR APPLICATION DATA:  (A) APPLICATION NUMBER: GB 9605656.9  (B) FILING DATE: 18-MAR-1996	
40	(2) INFORMATION FOR SEQ ID NO: 1:	
45	(i) SEQUENCE CHARACTERISTICS:  (A) LENGTH: 116 base pairs  (B) TYPE: nucleic acid  (C) STRANDEDNESS: single  (D) TOPOLOGY: linear	
	(ii) MOLECULE TYPE: cDNA to mRNA	
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(A) LENGTH: 1934 base pairs (B) TYPE: nucleic acid

(C) STRANDEDNESS: double

(D) TOPOLOGY: circular

(ii) MOLECULE TYPE: CDNA

(vi) ORIGINAL SOURCE:

(A) ORGANISM: Heliothis virescens

10

5

(vii) DAMEDIATE SOURCE: (B) CLONE: pSK19R

15

(xd) SEQUENCE DESCRIPTION: SEQ ID NO: 2:

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	(C) STRANDEDNESS: double (D) TOPOLOGY: circular	
	(ii) MOLECULE TYPE: cDNA	
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50	TCAGAAAACG AGTCAATGTC ATCAGGTCGT GAGGAACTGT CTCCAGCTTC GAGTGTAAAC	360
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60	TTGAAGAAAT GTCTTGCGGT GGGCATGAGG CCCGAGTGCG TGGTGCCGGA GAACCAGTGT	660

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10	OTTACACAGT	CGGACGAGGA	CGACGAAGAC	TCGGATATGC	COTTCCGTCA	GATTACCGAG	1020
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<sup>(2)</sup> INFORMATION FOR SEQ ID NO: 4:

(i)	SEQUI	ENCE CHARACTERISTICS:
		LENGTH: 2745 base pairs
	(B)	TYPE: nucleic acid
	(C)	STRANDEDNESS: doubl

(D) TOPOLOGY: linear

#### (ii) MOLECULE TYPE: CDNA

10 (ix) FEATURE:

(A) NAME/KEY: CDS

(B) LOCATION: 225..1955

(D) OTHER INFORMATION:/codon\_start= 225
/product= "Heliothis ecdysome receptor"

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	•				GACGACAGTA		1020
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	a manufa maam	AGAATTCGCT	A A CCCCCCTCC	CGGCTTCGC	CAAGATCTCG	CAGTCGGACC	1380
							1440
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	GTGATGAGTC	GTCCGCTGTC	CACGTCGCCG	TCACATGTTT	GTTTCTGATG	CACACGTGAG	2520
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45	AAGTGTGTCT	TATTACAATA	CAAAGTGTGT	GTCGTCGATA	GCTTCCACAC	GAGCAAGCCT	2700
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(2) INFORMATION FOR SEQ ID NO: 5:

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- (i) SEQUENCE CHARACTERISTICS:
  - (A) LENGTH: 575 amino acids
    (B) TYPE: amino acid
    (C) STRANDEDNESS: single
    (D) TOPOLOGY: linear
- 55
  - (ii) MOLECULE TYPE: protein

60

(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 5:

	Met 1	Ser	Leu	Gly	Ala 5	Arg	Gly	Tyr	Arg	Arg 10	CAR	Asp	Thr	Leu	Ala 15	Asp
<b>5</b>	Met	Arg	Arg	Arg 20	Trp	Tyr	Asn	Asn	Gly 25	Gly	Phe	Gln	Thr	Leu 30	Arg	<b>Net</b>
	Leu	Glu	Glu 35	Ser	Ser	Ser	Glu	Val 40	Thr	Ser	Ser	Ser	Ala 45	Leu	Gly	Leu
10	Pro	Pro 50	Ala	Met	Val	Met	Ser 55	Pro	Glu	Ser	Leu	Ala 60	Ser	Pro	Glu	Ile
15	Gly 65	Gly	Leu	Glu	Leu	71P	Gly	Tyr	Asp	ХSP	Gly 75	Ile	Thr	Tyr	Ser	Met 80
	Ala	Gln	Ser	Leu	Gly 85	Thr	СЛВ	Thr	Met	Glu 90	Gln	Gln	Gln	Pro	Gln 95	Pro
20	Gln	Gln	Gln	Pro 100	Gln	Gln	Thr	Gln	Pro 105	Leu	Pro	Ser	Met	Pro 110	Leu	Pro
	Met	Pro	Pro 115	Thr	Thr	Pro	Lys	Ser 120	Glu	Asn	Glu	Ser	Met 125	Ser	Ser	Gly
25	Arg	Glu 130	Glu	Leu	Ser	Pro	Ala 135	Ser	Ser	Val	Asn	Gly 140	СУВ	Ser	Thr	λsp
30	Gly 145	Glu	Ala	Arg	Arg	Gln 150	Lys	Lys	Gly	Pro	Ala 155	Pro	Arg	Gln	Gln	Glu 160
	<b>Gl</b> u	Leu	Cys	Leu	Val 165	СЛВ	Gly	Asp	Arg	Ala 170	Ser	Gly	Tyr	His	Tyr 175	Asn
35	Ala	Leu	Thr	Cys 180	Glu	Gly	Cys	Lys	Gly 185	Phe	Phe	Arg	Arg	Ser 190	Val	Thr
	Lys	λsn	Ala 195	Val	Tyr	Ile	Сув	Lys 200	Phe	Gly	His	Ala	Cys 205	Glu	Met	ДSP
40	Ile	Туг 210	Met	Arg	Arg	Lys	Cys 215	Gln	Glu	Сув	Arg	Leu 220	Lys	Lys	Сув	Leu
45	Ala 225	Val	Gly	Met	Arg	Pro 230	Glu	Сув	Val	Val	Pro 235	Glu	Asn	Gln	Cys	Ala 240
	Met	Lys	Arg	Lys	Glu 245	Lys	Lys	Ala	Gln	Arg 250	Glu	Lys	yab	Lys	Leu 255	Lio
50	Val	Ser	Thr	Thr 260	Thr	Val	yab	Хsр	His 265	Het	Pro	Pro	Ile	Met 270	Gln	Cys
	Asp		Pro 275	Pro	Pro	Glu	Ala	Ala 280	Arg	Ile	Leu	Glu	Сув 285	Val	Gln	His
55	Glu	Val 290	Val	Pro	Arg	Phe	Leu 295	Asn	Glu	Lys	Leu	<b>Met</b> 300	Glu	Gln	Asn	Arg
60	Leu 305	Lys	<b>As</b> n	Val		Pro 310	Leu	Thr	Ala	Asn	Gln 315	Lys	Ser	Leu		<b>Ala</b> 320
	Arg	Leu	Val		Tyr 325	Gln	Glu	Gly	Tyr	Glu 330	Gln	Pro	Ser		Glu 335	Asp

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		Leu	Lys	Arg	Val 340		Gln	Ser	Asp	Glu 345		Asp	Glu	Asp	Ser 350		Met
5		Pro	Phe	Arg 355		Ile	Thr	Glu	<b>Met</b> 360		11	Leu	Thr	Val 365	Gln	Leu	Ile
		Val	G1u 370		Ala	Lys	Gly	Leu 375		Gly	Phe	Ala	Lys 380		Ser	Gln	Ser
10		Asp 385	Gln	Ile	Thr	Leu	Leu 390	Lys	Ala	Сув	Ser	Ser 395	Glu	Val	Met	Met	Leu 400
15		Arg	Val	λla	Arg	Arg 405	Tyr	Asp	Ala	Ala	Thr 410	Asp	Ser	Val	Leu	Phe 415	Ala
		Asn	Asn	Gln	Ala 420	Tyr	Thr	Arg	Asp	Asn 425	Tyr	Arg	Lys	Ala	Gly 430	Met	Ala
20		Tyr	Val	Ile 435	Glu	Asp	Leu	Leu	His 440	Phe	Сув	Arg	Сув	Met 445	Tyr	Ser	Met
		Met	Met 450	Asp	Asn	Val	His	Tyr 455	Ala	Leu	Leu	Thr	Ala 460	Ile	Val	Ile	Phe
25		Ser 465	Asp	Arg	Pro	Gly	Leu 470	Glu	Gln	Pro	Leu	Leu 475	Val	Glu	Asp	Ile	Gln 480
30		Arg	Tyr	Tyr	Leu	Asn 485	Thr	Leu	Arg	Val	Tyr 490	Ile	Leu	Asn	Gln	<b>As</b> n 495	Ser
		Ala	Ser	Pro	Arg 500	Gly	Ala	Val	Ile	Phe 505	Gly	Glu	Ile	Leu	Gly 510	Ile	Leu
35		Thr	Glu	11e 515		Thr	Leu	Gly	Met 520	Gln	Asn	Ser	Asn	Met 525	СЛв	Ile	Ser
		Leu	Lys 530	Leu	Lys	Lys	Arg	Lys 535		Pro	Pro	Phe	Leu 540	Glu	Glu	Ile	Trp
40		λsp 545	Val	Ala	Asp	Val	<b>Ala</b> 550	Thr	Thr	Ala		Pro 555	Val	Ala	Ala	Glu	<b>Ala</b> 560
45		Pro	Ala	Pro	Leu	Ala 565	Pro	Ala	Pro	Pro	<b>Ala</b> 570	Arg	Pro	Ala	Thr	Val 575	-
	(2)	INFOR	MATI	CORF E	POR 'S	EQ 1	D NC	): 6:	٠								
<b>5</b> 0		. (1)	(A) (B) (C)	LENCI TYI STI TOI	GTH: PE: D VANDE	948 ucle	bas ic a S: d	e pa cid loubl	irs								
55		(ii) (vi)	ORIG		, sou	RCE:		pter	a ex	igua	ı						

(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 6:

	CANAGGGAAA AAGACAAGTT GCCAGTCAGT ACAACGACAG TGGATGATCA CATGCCTCCC	120
_	ATTATGCAGT GTGATCCACC GCCTCCAGAG GCCGCAAGAA TTCACGAGGT GGTGCCACGA	180
5	TTCCTGAATG AAAAGCTAAT GGACAGGACA AGGCTCAAGA ATGTGCCCCC TCACTGCCAA	240
	CCAGAAGTCC TTAATAGCGA GGCTGGTCTG GTACCAAGAA GGCTATGAAC AGCCATCAGA	300
10	AGAGGATCTA AAAAGAGTCA CACAGTCGGA TGAAGACGAA GAAGAGTCGG ACATGCCGTT	360
	CCGTCAGATC ACCGAGATGA CGATCCTCAC AGTGCAGCTC ATTGTTGAAT TCGCTAAGGG	420
	CCTACCAGCG TTCGCAAAGA TCTCACAGTC GGATCAGATC ACATTATTAA AGGCCTGTTC	480
15	GAGTGAGGTG ATGATGTTGC GAGTAGCTCG GCGGTACGAC GCGGCGACAG ACAGCGTGTT	540
	OTTCOCCAAC AACCAGGCGT ACACCCGCGA CAACTACCGC AAGGCAGGCA TGGCCTACGT	600
20	CATCGAGGAC CTGCTGCACT TCTGCCGGTG CATGTACTCC ATGATGATGG ATAACGTCCA	660
	CTATGCACTG CTCACTGCCA TCGTCATTTT CTCAGACCGA CCCGGGCTTG AGCTAACCCT	720
25	GTTGGTGGAG GAGATCCAGA GATATTACCT GAACACGCTG CGGGTGTACA TCCTGAACCA	780
ຜ	GAACAGTCGG TCGCCGTGCT GCCCTGTCAT CTACGCTAAG ATCCTCGGCA TCCTGACGGA	840
	GCTGCGGACC CTGGGCATGC AGAACTCCAA CATGTGCATC TCACTCAAGC TGAAGAACAG	900
30	GAACGTGCCG CCGTTCTTCG AGGATATCTG GGACGTCCTC GAGTAAAA	948
	(2) INFORMATION FOR SEQ ID NO: 7:	
35	(i) SEQUENCE CHARACTERISTICS:  (A) LENGTH: 319 amino acids  (B) TYPE: amino acid  (C) STRANDEDNESS: single  (D) TOPOLOGY: linear	
40	(ii) MOLECULE TYPE: protein	
45		
	(xi) Sequence Description: Seq ID NO: 7:	
	(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 7:  Arg Pro Glu Cys Val Val Pro Glu Asn Gln Cys Ala Net Lys Arg Lys	
	Arg Pro Glu Cys Val Val Pro Glu Asn Gln Cys Ala Met Lys Arg Lys 1 5 10 15	
<b>5</b> 0	Arg Pro Glu Cys Val Val Pro Glu Asn Gln Cys Ala Met Lys Arg Lys	
50	Arg Pro Glu Cys Val Val Pro Glu Asn Gln Cys Ala Met Lys Arg Lys  1 5 10 15  Glu Lys Lys Ala Gln Arg Glu Lys Asp Lys Leu Pro Val Ser Thr Thr 20 25 30  Thr Val Asp Asp His Met Pro Pro Ile Met Gln Cys Asp Pro Pro Pro	
50 55	Arg Pro Glu Cys Val Val Pro Glu Asn Gln Cys Ala Met Lys Arg Lys 1 5 10 15  Glu Lys Lys Ala Gln Arg Glu Lys Asp Lys Leu Pro Val Ser Thr Thr 20 25 30  Thr Val Asp Asp His Met Pro Pro Ile Met Gln Cys Asp Pro Pro Pro A0 45	
	Arg Pro Glu Cys Val Val Pro Glu Asn Gln Cys Ala Met Lys Arg Lys  1 5 10 15  Glu Lys Lys Ala Gln Arg Glu Lys Asp Lys Leu Pro Val Ser Thr Thr 20 25 30  Thr Val Asp Asp His Met Pro Pro Ile Met Gln Cys Asp Pro Pro Pro	
	Arg Pro Glu Cys Val Val Pro Glu Asn Gln Cys Ala Met Lys Arg Lys  Glu Lys Lys Ala Gln Arg Glu Lys Asp Lys Leu Pro Val Ser Thr Thr  20  Thr Val Asp Asp His Met Pro Pro Ile Met Gln Cys Asp Pro Pro Pro  35  Pro Glu Ala Ala Arg Ile Leu Glu Cys Val Gln His Glu Val Val Pro	

	Tyr	ln	Glu	Gly 100	Tyr	Glu	Gln	Pro	Ser 105	Glu	Glu	Asp	Leu	Lys 110	λrg	Val
5	Thr	Gln	Ser 115	<b>As</b> p	Glu	λsp	λsp	Glu 120	λsp	Ser	Авр	Met	Pro 125	Phe	λrg	Gln
10	Ile	Thr 130	Glu	Met	Thr	Ile	Leu 135	Thr	Val	Gln	Leu	Ile 140	Val	Glu	Phe	Ala
	Lys 145	Gly	Leu	Pro	Gly	Phe 150	Ala	Lys	Ile	Ser	Gln 155	Ser	Asp	Gln	Ile	Thr 160
15	Leu	Leu	Lys	Ala	Cys 165	Ser	Ser	Glu	Val	<b>Met</b> 170	Met	Leu	Arg	Val	Ala 175	λrg
	Arg	Tyr	Aep	Ala 180	Ala	Thr	Asp	Ser	<b>Val</b> 185	Leu	Phe	Ala	Asn	<b>Asn</b> 190	Gln	Ala
20 .	Tyr	Thr	Arg 195	Asp	Asn	Tyr	Arg	Lys 200	Ala	Gly	Met	Ala	Tyr 205	Val	Ile	Glu
25	Asp	Leu 210	Leu	His	Phe	Сув	Arg 215	Сув	Met	Tyr	Ser	Met 220	Met	Met	Asp	Asn
_	Val 225	His	Tyr	Ala	Leu	Leu 230	Thr	Ala	Ile	Val	Ile 235	Phe	Ser	Asp	Arg	Pro 240
30	Gly	Leu	Glu	Gln	Pro 245	Leu	Leu	Val	Glu	Glu 250	Ile	Gln	Arg	Tyr	Tyr 255	Leu
	Asn	Thr	Leu	<b>Arg</b> 260	Val	Tyr	Ile	Leu	Asn 265	Gln	Asn	Ser-	Ala	Ser 270	Pro	Arg
35	Gly	Ala	Val 275	Ile	Phe	Gly	Glu	11e 280	Leu	Gly	Ile	Leu	Thr 285	Glu	Ile	Arg
40	Thr	Leu 290	Gly	Met	Gln	Asn	Ser 295	Asn	Met	Сув	Ile	Ser 300	Leu	Lys	Leu	Lys
	Lys 305	Arg	Lys	Leu	Pro	Pro 310	Phe	Leu	Glu	Glu	Ile 315	Asp	Trp	Asp	Val	

#### **CLAIMS**

- 1. DNA comprising the sequence shown in Seq ID No. 2.
- 5 2. DNA comprising the sequence shown in Seq ID No. 3.
  - 3. DNA comprising the sequence shown in Seq ID No. 4.
- 4. DNA comprising a sequence which shows 60% or more homology with the sequence shown in Seq ID No 1, 2 or 3.
  - 5. DNA according to claim 4 wherein said homology is in the range of 65% to 99%.
- 6. DNA which hybridises to the sequence shown in Seq. ID No. 2, 3 or 4, and which codes for at least part of the *Heliothis* ecdysone receptor.
  - 7. DNA which is degenerate as a result of the genetic code to the DNA of any one of claims 1 to 6 and which codes for a polypeptide which is at least part of the *Heliothis* ecdysone receptor.

8. DNA comprising part of the sequence shown in Seq ID No. 2, and which codes for at least part of the *Heliothis* ecdysone receptor ligand binding domain.

- 9. DNA comprising part of the sequence shown in Seq ID No. 3, and which codes for at least part of the *Heliothis* ecdysone receptor ligand binding domain.
  - 10. DNA comprising part of the sequence shown in Seq ID No. 4, and which codes for at least part of the *Heliothis* ecdysone receptor ligand binding domain.
- DNA comprising a sequence which shows 60% or more homology with the sequence of claim 8, 9 or 10.
  - 12. DNA according to claim 11 wherein said homology is in the range of 65% to 99%.
- 35 13. DNA which hybridises to the DNA of any one of claims 8 to 12 and which codes for at least part of the Heliothis ecdysone receptor ligand binding domain.

- 14. DNA which is degenerate as a result of the genetic code to the DNA of any one of claims 8 to 12 and which codes for a polypeptide which is at least part of the *Heliothis* ecdysone receptor ligand binding domain.
- 5 15. DNA comprising part of the sequence shown in Seq ID No. 2, and which codes for at least part of the *Heliothis* ecdysone receptor DNA binding domain.
  - 16. DNA comprising part of the sequence shown in Seq ID No. 3, and which codes for at least part of the *Heliothis* ecdysone receptor DNA binding domain.
  - 17. DNA comprising part of the sequence shown in Seq ID No. 4, and which codes for at least part of the *Heliothis* ecdysone receptor DNA binding domain.
- 18. DNA comprising a sequence which shows 60% or more homology with the sequence of claim 15, 16 or 17.
  - 19. DNA according to claim 18 wherein said homology is in the range of 65% to 99%.
- 20. DNA which hybridises to the DNA of any one of claims 15 to 19 and which codes for
   at least part of the Heliothis ecdysone receptor DNA binding domain.
  - 21. DNA which is degenerate as a result of the genetic code to the DNA of any one of claims 15 to 19 and which codes for a polypeptide which is at least part of the Heliothis ecdysone receptor DNA binding domain.
- 22. DNA comprising part of the sequence shown in Seq ID No. 2, and which codes for at least part of the *Heliothis* ecdysone receptor transactivation domain.
- DNA comprising part of the sequence shown in Seq ID No. 3, and which codes for at
   least part of the Heliothis ecdysone receptor transactivation domain.
  - 24. DNA comprising part of the sequence shown in Seq ID No. 4, and which codes for at least part of the *Heliothis* ecdysone receptor transactivation domain.
- DNA comprising a sequence which shows 60% or more homology with the sequence of claim 22, 23 or 24.

- 26. DNA according to claim 25 wherein said homology is in the range of 65% to 99%.
- 27. DNA which hybridises to the DNA of any one of claims 22 to 26 and which codes for at least part of the *Heliothis* ecdysone receptor transactivation domain.
- 28. DNA which is degenerate as a result of the genetic code to the DNA of any one of claims 22 to 26 and which codes for a polypeptide which is at least part of the Heliothis ecdysone receptor transactivation domain.
- DNA comprising part of the sequence shown in Seq ID No. 2, and which codes for at least part of the *Heliothis* ecdysone receptor hinge domain.
  - 30. DNA comprising part of the sequence shown in Seq ID No. 3, and which codes for at least part of the *Heliothis* ecdysone receptor hinge domain.
- DNA comprising part of the sequence shown in Seq ID No. 4, and which codes for at least part of the Heliothis ecdysone receptor hinge domain.
- DNA comprising a sequence which shows 60% or more homology with the sequence of claim 29, 30 or 31.
  - 33. DNA according to claim 32 wherein said homology is in the range of 65% to 99%.
- 34. DNA which hybridises to the DNA of any one of claims 29 to 33 and which codes for at least part of the *Heliothis* ecdysone receptor hinge domain.
  - 35. DNA which is degenerate as a result of the genetic code of the DNA of any one of claims 29 to 33 and which codes for a polypeptide which is at least part of the *Heliothis* ecdysone receptor hinge domain.
  - 36. DNA having part of the sequence shown in Seq ID No. 2, and which codes for at least part of the *Heliothis* ecdysone receptor carboxy terminal region.
- DNA having part of the sequence shown in Seq ID No. 3, and which codes for at least
   part of the Heliothis ecdysone receptor carboxy terminal region.

- 38. DNA having part of the sequence shown in Seq ID No. 4, and which codes for at least part of the Heliothis ecdysone receptor carboxy terminal region.
- 39. DNA comprising a sequence which shows 60% or more homology with the sequence of claim 36, 37 or 38.
  - 40. DNA according to claim 39 wherein said homology is in the range of 65% to 99%.
- DNA which hybridises to the DNA of any one of claims 36 to 40 and which codes for at least part of the Heliothis ecdysone receptor carboxy terminal region.
  - DNA which is degenerate as a result of the genetic code of the DNA of any one of claims 36 to 40 and which codes for a polypeptide which is at least part of the Heliothis ecdysone receptor carboxy terminal region.
- 43. A polypeptide comprising the Heliothis ecdysone receptor or a fragment thereof, wherein said polypeptide is substantially free from other proteins with which it is ordinarily associated, and which is coded for by the DNA of any preceding claim.
- 20 44. A polypeptide comprising the amino acid sequence shown in Seq ID No. 4 or any allelic variant or derivative thereof.
- A polypeptide comprising part of the amino acid sequence shown in Seq ID No. 4 or any allelic variant or derivative thereof, which sequence provides the *Heliothis* ecdysone receptor ligand binding domain.
  - 46. A polypeptide comprising part of the amino acid sequence shown in Seq ID No. 4 or any allelic variant or derivative thereof, which sequence provides the *Heliothis* ecdysone receptor DNA binding domain.
- 47. A polypeptide comprising part of the amino acid sequence shown in Seq ID No. 4 or any allelic variant or derivative thereof, which sequence provides the Heliothis ecdysone receptor transactivation domain.
- A polypeptide comprising part of the amino acid sequence shown in Seq ID No. 4 or any allelic variant or derivative thereof, which sequence provides the *Heliothis* ecdysone receptor hinge domain.

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- 49. A polypeptide comprising part of the amino acid sequence shown in Seq ID No. 4 or any allelic variant or derivative thereof, which sequence provides the *Heliothis* ecdysone receptor carboxy terminal region.
- 50. A polypeptide according to any one of claims 44 to 49 wherein said derivative is a homologous variant which includes conservative amino acid changes.
- 51. DNA comprising the sequence shown in Seq ID No. 6.
- 52. DNA comprising a sequence which shows 60% or more homology with the sequence shown in Seq ID No. 6.
- 53. DNA according to claim 52 wherein said homology is in the range of 65% to 99%.
- 54. DNA which hybridises to the DNA sequence shown in Seq ID No. 6 and which codes for at least part of *Spodoptera* ecdysone receptor.
- 55. DNA which is degenerate as a result of the genetic code to the DNA of any one of claims 51 to 54.
  - 56. DNA comprising part of the sequence shown in Seq ID No. 6, and which codes for at least part of the Spodoptera ecdysone receptor ligand binding domain.
- 25 57. DNA comprising a sequence which shows 60% or more homology with the sequence of claim 56.
  - 58. DNA according to claim 57 wherein said homology is in the range of 65% to 99%.
- 30 59. DNA which hybridises to the DNA of any one of claims 56 to 58 and which codes for at least part of the Spodoptera ecdysone receptor ligand binding domain.
- 60. DNA which is degenerate as a result of the genetic code to the DNA of any one of claims 56 to 58 and which codes for at least part of the Spodoptera ecdysone receptor
   35 ligand binding domain.

- 61. DNA comprising part of the sequence shown in Seq ID No. 6, and which codes for at least part of the Spodoptera ecdysone receptor hinge domain.
- 62. DNA comprising a sequence which shows 60% or more homology with the sequence of claim 61.
  - 63. DNA according to claim 62 wherein said homology is in the range of 65% to 99%.
- 64. DNA which hybridises to the DNA of any one of claims 61 to 63 and which codes for at least part of the Spodoptera ecdysone receptor hinge domain.
  - 65. DNA which is degenerate as a result of the genetic code to the DNA of any one of claims 61 to 63 and which codes for at least part of the *Spodoptera* ecdysone receptor hinge domain.
  - 66. A polypeptide coded for by the DNA of any one of claims 51 to 65.
  - 67. A fusion polypeptide comprising the polypeptide of claim 45 or 50 (when dependent upon claim 45) and functionally linked to a DNA binding domain and a transactivation domain.
    - 68. Recombinant DNA comprising the DNA of any one of claim 8 to 14 functionally linked to DNA encoding a DNA binding domain and a transactivation domain.
- 25 69. A fusion polypeptide according to claim 67 or recombinant DNA according to claim 68 wherein the DNA binding domain and/or transactivation domain is fungal, bacterial, plant or mammalian.
- 70. A fusion polypeptide or recombinant DNA according to claim 69 wherein the DNA binding domain is GALA or A1CR/A.
  - 71. A fusion polypeptide or recombinant DNA according to claim 69 or 70 wherein the transactivation domain is VP16.
- 35 72. A fusion polypeptide or recombinant DNA according to claim 69 wherein the DNA binding domain and/or transactivation domain is from a steroid receptor superfamily member.

- 73. A fusion polypeptide or recombinant DNA according to claim 72 wherein the DNA binding domain and/or transactivation domain is from a glucocorticoid or a Spodoptera ecdysone receptor.
- 74. A recombinant DNA construct comprising recombinant DNA of any one of claims 68 to 73; and DNA which codes for a gene operably linked to a promoter sequence and a hormone response element, which is responsive to the DNA binding domain coded for by said recombinant DNA.
- 75. A fusion polypeptide comprising the polypeptide of claim 46 or 50 (when dependent upon claim 46) and functionally linked to a ligand binding domain and a transactivation domain.
- 15 76. Recombinant DNA comprising the DNA of any of claims 15 to 21 functionally linked to DNA encoding a ligand binding domain and a transactivation domain.
  - 77. A fusion polypeptide according to claim 75 or recombinant DNA according to claim 76 wherein the ligand binding domain and/or transactivation domain is fungal, bacterial, plant or mammalian.
  - 78. A fusion polypeptide or recombinant DNA according to claim 77 wherein the transactivation domain is VP16.
- 25 79. A fusion polypeptide or recombinant DNA according to claim 77 wherein the ligand binding domain and/or transactivation domain is from a steroid receptor superfamily member.
- 80. A fusion polypeptide or recombinant DNA according to claim 79 wherein the ligand binding domain and/or transactivation domain is from a glucocorticoid or Spodoptera ecdysone receptor.
- 81. A recombinant DNA construct comprising recombinant DNA of any one of claims 76 to 80; and DNA which codes for a gene operably linked to a promoter sequence and a hormone response element, which is responsive to the DNA binding domain coded for by said recombinant DNA.

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- 82. A fusion polypeptide comprising the polypeptide of claim 47 or 50 (when dependent upon claim 47) and functionally linked to a ligand binding domain and a DNA binding domain.
- 83. Recombinant DNA comprising the DNA of any one of claims 22 to 28 functionally linked to DNA encoding a ligand binding domain and a DNA binding domain.
  - 84. A fusion polypeptide according to claim 82 or recombinant DNA according to claim
     83 wherein the ligand binding domain and/or DNA binding domain is fungal, bacterial,
     plant or mammalian.
    - 85. A fusion polypeptide or recombinant DNA according to claim 84 wherein the DNA binding domain is GAL4 or A1CR/A.
- 15 86. A fusion polypeptide or recombinant DNA according to claim 84 wherein the ligand binding domain and/or DNA binding domain is from a steroid receptor superfamily member.
- 87. A fusion polypeptide or recombinant DNA according to claim 86 wherein the ligand binding domain and/or DNA binding domain is from a glucocorticoid or *Spodoptera* ecdysone receptor.
- 88. A recombinant DNA construct comprising recombinant DNA of any one of claims 82 to 87; and DNA which codes for a gene operably linked to a promoter sequence and a hormone response element, which is responsive to the DNA binding domain coded for by said recombinant DNA.
  - 89. A recombinant DNA construct comprising DNA according to any one of claims 1 to 7; and DNA comprising a sequence which codes for a gene operably linked to a promoter sequence and at least one hormone response element which is responsive to the DNA binding domain coded for by said DNA of any one of claim 1 to 7.
  - 90. A recombinant DNA construct according to any one of claims 74, 81, 88 and 89 wherein said promoter sequence codes for a constitutive, spatially or temporally regulating promoter.

- 91. A recombinant DNA construct according to any one of claims 74, 81, 88 and 89 wherein there is more than one copy of the hormone response element.
- 92. A cell transformed with the DNA of any one of claims 1 to 42, and 51 to 65; the polypeptide of any one of claims 43 to 50; the fusion polypeptide of any one of claims 67, 70 to 73, 75, 77 to 80, 82 and 84 to 87; the recombinant nucleic acid of any one of claims 68 to 73, 76 to 80 and 85 to 87; or the recombinant DNA construct of any one of claims 74, 81, 88 and 89.
- 10 93. A cell according to claim 92 wherein said cell is a plant, fungal or mammalian cell.
  - 94. A plant, fungus or mammal comprising the recombinant DNA construct of any one of claims 74, 81, 88 and 89.
- A method of selecting compounds capable of being bound to an insect steroid receptor superfamily member comprising screening compounds for binding to said polypeptide of any one of claims 43 to 50 or the fusion polypeptide of any one of claims 67, 70 to 73, 75, 77 to 80, 82 and 84 to 87, and selecting said compounds exhibiting said binding.
- 2096. A compound selected using the method of claim 95.
  - 97. An agricultural or pharmaceutical composition comprising the compound of claim 96.
- 25 98. Use of the compound of claim 96 as an agrochemical or a pharmaceutical.
  - 99. A method of producing a protein, peptide or polypeptide comprising introducing into the cell of claim 92, a compound which binds to the ligand binding domain in said cell.

Fig. 1.

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Sequence ID

TGCG AGG GGT GCA AGG AGT TCT TCA GGC GGA GTG TAA CCA AAA ATG TTT TAC ACGC TCC CCA CGT TCC TCA AGA AGT CCG CCT CAC ATT GGT

GAA CGC TTT ACC TAT ACA CAG TGT ACA TAT GCA AAT TCG GCC ATG CTT GCG AAA TGG ATA TGT ACA TGT ATA CGT TTA AGC CGG TAC GIC

1/56

ATA TGC GGA GAA AAT GCC AAG AGT A
TAT ACG CCT CTT TTA CGG TTC TCA T

91

SUBSTITUTE SHEET (RULE 26)

AGC

GGA GGA

SCA CGT

ည် ကို

00 CA

777 766

AAT

ဗ္ဗ ဗ္ဗ ဗ္ဗ

ပ္ပပ္ပ ဗိမ္မ

TTC

TCA

ATA

AGA

316

CTC

GTG

GTG

000 000 000

ATG

GTG

**000** 

TGT

AAC

GTA

CTG

CCT

GAC

ATG

166 ACC

7
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귱

45	TTA	CTG	CCT	TCG	AAG	CAG
	CAT	၁၁၁	ACC	990 000	TTG	AGA
39	GG T	CCA	ATG	ACT	TCA	AGA
	TCT	TGC	AGT	CCA GGT	AGA	ပ္ပပ္ပ (၁)
33	- ၁၁၅ ၁၁၅	TGC ACG	AGC TCG	CAA	TCG	CCA
	AAG	TCC	TGA	GCA	ACT	CTG
27	GAA	ATC	CCT GGA	CCT	ACA	SGT GGT
	ACA TGT	ATC	CAA	CCT	atg	CCA
21	ACC TGG	GTC CAG	TGT	ACG TGC	TCC	ACG
	ACC TGG	AAG	13GG ACC	CCA	TCA	TAC
15	TTC	AAG	CGT	TCT	AGG	CTG
	GTT	GCT	CGT	TCA	CTA	CCA GGT
თ —	GGT CCA	GGT CCA	GTT	AGG	TCG	TGA
	ACT TGA	GGT CCA	CAT	ACA	CTC	TGA
<u>ن</u> –	AGG	GAG	<b>A</b> CC <b>1</b> 66	CTT	CTC	CAC CAC
	-	46	91	136	181	226

បថ	ဗ ပ	4 F	4 E	A H	ធូ ប្	ပ္ဟ ပ္ပ	TCG	ES	GCA
ည္သတ္သ	TCG AGC	AGA	S C C C	TCA	ဗ္ဗ ဗ္ဗ	TGC	- •	GGT CCA	
AAT	AGG TCC	CAC	GC CGT	ATA TAT	CAG	CCA	GTG	CGA	999
ACC TGG	ATC	CAG	GAG	၁ ၁ ၁	CTT	ဗ္ဗဗ္ဗ ဗဗ္ဗဗ	GGA	GTG	AAA
GTT	GTC	CTG	ပ္ပ ဗ္ဗ	CTC	TTT	ATT	TCA	CGA	GAA
ပ္ပဗ္ဗ	AAT	၁၂၂	AGC	AGC	AGG	CAA	ATG	000 000	AGA
CAT	GTC	AAA T'T'T	ပ္ပပ္ပ ဗဗ္ဗ	CAG	TAA	ATG	AAA	GAG	GAA
TTC	CGA	TGT	AGG	CGA	GTG	CAT	GAG	CAT	ACG
<b>A</b> CC 1766	AAA	GAG	GRA	၁၁၁	AGG	GTA	၁၁၁ ၁၅၁	000 000	GAA
CCT	AGA	TTC	GAA	CTG	TGA	AGT	TAT	GGT	AAT
<b>A</b> CC <b>1</b> 66	ATC	AGC	GCA	TGT	ATG	TGC	CTA	TGC	TGC
ACA	CAA	TCC	<del>၀</del> ၀၀ ၁၅၁	TCT	CAC	AAA TTT	TAT	TCT	GTG
GAC	ACC	GTC	GAG	ATG	GCT	CAA	GGA	ATG	CCA GGT
GCA	AAC	ACT	၁၅၅	GGA CGA	၁၅၁ ၁၅၁	AAC	AAT TTA	GAA	GAA
GCA CGT	GAC	GGA	CGA	AGA	GTT	TGT	CGA	CTT	GGA
CCT	ACC TGG	TGA	TGG	AGA	CTA	GAG	TTG	GTT	ည် ပို့ပို့
361	406	451	496	541	586	631	929	721	766
									•

GAT GTG GAG S S GTC ACT TGA ATA TRA 000 S S SCT S 000 ပ္ပုပ္ပ ဗွ CAT GCA CGT GAT g g CGT GCA GAR हु सु AGA AGA 300 TAT GA CTC GG4 950 CGA CGA CCT GGA g CGA AGT 5 ည် စိ ပ္ပ ဗွ GGA CCT 95 GAT TGA ACT GCT AAG AGA TCT 350 SE မှ မှ မှ SCT S ATT ပ္ပ်ပ္ပ ပ္ပပ္ပ CTC AGT CAR AAG CAC GAC 700 A S S S CAC GIG ပ္ပ်ပ္ပ ပိပ္ပဲ CTC 000 GAT S S S S AGA TAC **CAG STC** ပ္ပ် ဗွ S C C TCT AGA CIT 91G CAC ပ္ပ ဗ္ဗ ဗ္ဗ CAG GAA GIT TTC NAG GC GC TGA ACT 5 TGA GAT 999 GTC ACC TGG CGA ည် သို့ S S S S S ATG S S S S S ACA TGT GTA GGA CCT GAA ACA GGA GAC 000 000 AAA **8**69 ပ္ပ ဗွ TGT GAT CCT ATT ဗ္ဗ ဗ္ဗ ဗ္ဗ TGA CGA GCT ATA TAA ATG TAT AGA AGA CAT 70g GGP PCP ဗ္ဗဗ္ဗ ပ္ပ CAA CTA GGA GIT 36 GCT Gra SC T 0 0 0 0 0 0 0 0 GTC TAC TAA CAC AGA TCT GAA 7GC ACG 700 TCT 770 A GAT SGA CCT ACA ပ္ပ ပ္ပ ပ္ပ GAT AGT A E ည် တို့ AAT TGA ACT CAC TAC ATT SG T 900 S F TCA GAA CCT S CT SS CAT TAG GTA S S **9**00 TCT S C GG P ည် အ GAG CTC A P g g 1126 1216 1261 811 946 991 1036 1081 1171 856 901 Fig.2 ii.

ပ္တတ္တ	GTG	TAC	CCT	GGT	CAT	999 CCC	CAG	CGT GCA	TCT
GAR	TCG	GCT	ACC	ACG	CGT	GCT	GAA	GGA CCT	0 0 0 0 0 0
ဗ္ဗ ဗ္ဗ ဗ္ဗ	CTG	CG.T	GG.	GCT	ပ္ပင္ပ ဗင္လပ္	CAC GTG	GAA	ပ္တ ပ္ပ	ပ္ပ ပ္ပ
CTA	GAA	TGC	TGA	CAC GTG	ဗ္ဗာ ဗ္ဗာ ဗ္	9 9 9 9	GCT	CGT	0 0 0 0 0
CAA	GGA CGT	TTA	GCT	SE C	ဗီပ ဗီပ ဗီ	GAT	CAA	GGA	ည် လူတွင်
CGA	GCT	55	0 0 0 0 0 0	CC CC CC CC	ပ္ပံ ပုပ္ပံ	GGA	CCT	CTG	ods CCT
TCG	CCT	CGT GCA	ပ္ပင္ပ ပုဇ္ပင္ပ	TTA	GTC	GAC	CTC	GAT	ပ္ပေပ
CAC	GGA	TAA	ပ္ပ်ပ္ ၁၅၅	ATA TAT	ပ္ပ ဗ္ဗ ဗ္ဗ	ACT	CAT	GGA	၁ ၁ ၁ ၁
GTA CAT	CGA	GGA	AGA	GAG	CAG	CAT	GTG	CGA	GGT CCA
ပ္ပပ္ပ ပပ္ပပ	CAT	GAT	CTC	CCA GGT	GAA	999 CCC	CAT	CCT	000
CCA	CGT	GAT	CTT	CAT	CCA	CCT	CAA	CAS	GAC
CAA	GTA	CAT	CAT	CCT	GAA	GAT	CTC	000	999
GAA	၁၅၁	CTC	TGT	GGA CCT	CCT	CGA	GAA	999	GAC
ပ္ပတ္ ဗပ္ပဗ္	CAT	GTA	CAT	CCA CCA	CAT	၁၁၁ ၁၁၁	GCA	GCT	GAC CTG
CAA	AGG	CAT	<b>3</b> 000	GAT	GTA	CTT	CAT	GRA	900
1306	1351	1396	1441	1486	1531	1576	1621	1666	1711

# Fig.2 iv

ပ္ပ ပ္ပ ပ	GCA CGT	CGA	<b>8</b> 8
ဗ္ဗ ဗ္ဗ ဗ	AGT TCA	CTG	TAC
GAT	TGA	GGA	10G
5 5	TAG	TAA	ATT
STG GTG	TTT	TTA	ပ္ပ ပ္ပ ပ္ပ ပ္ပ
ပ္ပပ္ပ	TAG	TAT	TAC
ပ္ပပ္ပ	999	ACC	ဗ္ဗ ဗ္ဗ ဗ္ဗ
ပ္ပ်ပ္ပ် ပုပ္ပံပဲ ပုပ္ပံပဲ	ACT	TCA	CAC
0 0 0 0 0	TAG	TGA	GCA CGT
ပ္ပ ပ္ပ ပ္ပ	TCA	ACG	<b>AGG</b>
ပ္ပ ဗ္ဗ	ည ၁၅၁၅	TCG	AAG
ပ္ပ် ပ္ပံ ပ္ပံ ပ္	GAA	ACG	CTT
ပ္ပ ပ္ပ	AGA	CTG	GCA GGT
ပ္ပဲ ပို့ ပို့	AGG	ACA	TTA
76c 76c	CTC	၁၁၁	ATT
1756	1801	1846	1891

Total number of bases is: 1934.

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The sequence shown below is that of pSK16.1

Sequence ID3

45	GAG	၅ ၂၂ ၂၂	ATC	<b>A</b> GC <b>1</b> 7CG	ပ္ပပ္ပ
	CTC CARC CARC C	CTG C	SAG 7	TAC 1	CAG (
39	ATG TAC G	0 000	222	ACT 1	CAG C
	CGA A	CTG G	17CG 0	ATC A	CAG C
33	•	SCA C	300 1 300 1	30C 1	and carc
	ACG O	rca g Agt c	CTC GAG G	GAT CTA C	ATG C
27		rcr 1	PCG O	GAC CTG	ACC 7
	AAG O	TCG 7	SAA	TAC (ATG (	TGC A
77	• •	FGC 7	ညည ညည	200	ACC
	GGA	GTG	TCC AGG	TGG	၁၉၁
15	AAC	GAG	ATG	CTG	CTG
	AAC	TCT	GTG	GAG	TCG
o —	TAT	TCG	atg Tac	CTG GAC	CAG
	700 ACC	AGC	GCT	ပ္ပံ ဗိပ္ပံ ဗိ	GCA CGT
m —	၁၀ ၁၀ ၁၀	GAG	ပ္ပပ္ပ ဗိဗ္ဗ	ပ္ပပ္ပ ပပ္ပပ္	ATG
	н	46	91	136	181

## Fig.3 .

TAC	TCA	AAC	CCA	AGA	AAA TTT	<b>TGC</b>
<b>300</b>	GAG	GTA	000 000	GAC	ACA	ATA
<b>6</b> 00	AAC	AGT	AAA	၅၁၁ ၁၅၅	000 000	TAC
GAT	CTT	TCG	AAG	TGC	GAA	GIG
9 9 9 9	TCA	GCT	CAG	GTC	TGT	Ş
GTT B	AATT	CCA	000 000	CTT	ACA	AAT
164 164	ပ္ပ်ပ္ ၁၁၁ ၁၅၅	TCT	AGG	TGT	CTC	ACC AAA
	ACA TGT	CTG	900	CTA	၁ ၁ ၁ ၁ ၁	
910	ACA TGT	GRA	GAG	GAG	AAC	GTA
ဗ္ဗ ဗ္ဗ	၅ ၂၀ ၂၀	GAG	၁၅ ၁၅ ၁၂	GAA	TAC	AGT
910 910	SGT GGT	CGT	GAT	CAA	CAC	S S S
61C	ATG	GGT CCA	ACA	CAG	TAT	AGG
	65 t	TCA	AGC TCG	AGG TCC	GGA CCT	TTC
9 <b>9</b>	TTA	TCA	TGC	ပ္သည္ ဗင္သည	TCC	TTC
	0 0 0 0 0	atg	00 00 00 00 00 00 00 00 00 00 00 00 00	ဗ္ဗာဗ ဗ္ဗာဗ	000 000	GGT
97	71	16	61	90	51	961

ACG	AAA	AGG	AAA	ACG	CCT	GAG	AGA	ATC	GAG	GAC CTG
TAT 1	AGA TCT	ATG TAC	993	AGT	GAC	CAC	AAC	TTG	TCC	S E
ATG	990	၁ဗ္ဗာ	AAA	OTC CAG	TGT	CAG	CAG GTC	10G	CCT SG A	GAC
CAC	ATG	GTG	ATG	ပ္ပပ္ပ ဗဗ္ဗ	CAA	GTG	GAA	AAG	GIT	GAC
CGT	TAT	999	S Corr	TTG	ATG	TGT	ATG	CAG	GAA	GE CO
TTA	ATC	GRA	TGT	AAA	ATC	GAA	CTA	aat Tta	TAT	GAC
TII	GAT	TGT	CAG	GAC	၁၁၁ ၁၁၁	CTG	AAG	ပ္ပပ္ပ ပုပ္ပပ္	၁၈ ၁၈ ၁၈	10G
<b>TGG</b>	ATG	AAA	AAC	AAA	CCT	ATT	GAG	ACT	GAA	CAG OTC
CAT	SE	AAG	GAG	GAA	ATG	AGA	AAT	CTC	CAG	ACA TGT
13	16C	TTG	000 000	AGG	CAC	GCT	CTG	000 000	TAC	F 2
သဗ	S S	ဗ္ဗ ဗ္ဗ ဗ္	GTG	CAG	GAT	ပ္ပ္ပ် ပ္ပပ္ပ	TTC	ပ္ပ <u>ဗ</u>	100 100	AGG 700
TCC	STATE OTTA	TGT ACA	GTG	900 000	GAC	GAG	CGA	GTG CAC	GTG	AAG
AAG	ပ္က ဗ္	GAG	TGC ACG	AAG	GTA	CCA GGT	CCA	AAC	CTC	CTG
AAG	TTC	CAG	GAG	AAT	ACA	000 000	GTG	AAG	AGG	GAC
CC	AATT	TGT	000 000	GAG	ACG TGC	၅ ၅ ၂၅	GTG	TTG	GCA CGT	GRA
	541	586	631	919	721	766	811	856	901	946

10/50

166	10g	GAT	ATG	ဗ္ဗ ပ္ပ ဗ္ဗ	TTC	g g	CAG	ATT	**************************************	QAG CTC	ATG	<b>30</b> 50 130 130 130 130 130 130 130 130 130 13	ATT	CTC GAG	ACA
1036	GTG CAC	CAG GTC	CTC	ATC	GTA CAT	GRA	TTC	SCT CGA	AAG	ပ္ပပ္ပ ပပ္ပပ္	CTC	900	999 000 000	TTC	ပ္တတ္တ
1081	AAG	atc	TCG AGC	CAG	TCG	GAC	CAG GTC	ATC	Acg TGC	TTA AAT	TTA AAT	AAG	ဗ္ဗ ဗ္ဗ ဗ္ဗ	10C	AGT.
1126	AGT	GAG	GTG	ATG	ATG	CTC GAG	S C T T	GTG	S S S S S	990 800	9 9 9	TAT ATA	GAC CATO	ဗ္ဗ ဗ္ဗ ဗ္ဗ	ပ္ပဗ္ဗ
1171	166 166	GAC	AGC	GTA	CTG	TTC	900	AAC	AAC	CAG	9 9 9 9	TAC	ACT	ပ္ပ ပ္ပ ပ္ပ	GAC
1216	AAC	TAC	000 000	AAG	GGT GGT	000 000	ATG	000 000	TAC	GTC	ATC	GAG	GAC	CTG	CTG
1261	CAC	TTC	TGT	ဗဗ္ဗ ဗဗ္ဗ	TGC	ATG	TAC	TCC	ATG	ATG	atg	GAT	AAC	GTG CAC	CAT
1306	TAT ATA	900	CTG	CIT	ACA	ပ္ပ ပ္ပဗ္ဗ	ATT	GAG CAG	atc	TTC	TCA	GAC	900	၁ ၁ ၁ ၁ ၁	ဗ္ဗ
1351	CTT	GAG CTC	CAA	ည ည	CTG	TTG	GTG	GAG	GAC	ATC	CAG	AGA	TAT	TAC	CTG
1396	AAC	ACG TGC	CTA	990	GTG	TAC	ATC	CITG GAC	AAC	CAG GTC	AAC	AGC	ဗ္ဗ ဗ္ဗ ဗ္ဗ	TCG AGC	ပ္ပ ဗွ
1441	ည္သည	ပ္ပစ္ပ	ညည	GTC	ATC	TTC	ည	GAG	ATC	CTC	ည	ATA	CTG	ACG	GAG

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SE SE SE SE SE SE SE SE SE SE SE SE SE S	CTC	166 ACC	SE CITO	ပ္တ ဗ္ဗ	AGT	ATT TAA	ACC TGG	STS CAC	ATT	PAT OF AT
<b>1</b> 3c	100 A	ATC	999	ည တ တ	GCT	CCT GGA	CG SC SC SC SC SC SC SC SC SC SC SC SC SC	CGT	TAT	ပ္ပတ္သ ပ္ပတ္သ ပ္ပတ္သ
GAC	ATC	GAG	ဗ္ဗ ဗ္ဗ ဗ္ဗ	900	CTG	CAA	ACC TGG	GAG	ATA TAT	ပ္ပ ၅ ဗွ ဗွ
TAT	13GC	GAG	GTG	900	AGA	GAT	CAC	GCA CGT	AGA	O STC
900	ATG	GAG	999	000	CAT	CGT	999	GAT	166 ACC	<del>ပ</del> ္ပံ ပ္ပံ
GAC	AAC	TTC	ACG 76C	000 000	GCT	CGA	AGA	GAC	ATT	70g 70gC
TAG	TCC AGG	၁၁၁	900	900	AAC	CGT	TTA AAT	GAC	ACG	ဗ္ဗ ဗ္ဗ
CIC	MAC	၁၁၁	ACG TGC	၁ ၁ ၁ ၁ ၁ ၁	GAG	TGA	CAC	ACC	TGA	16G
99	0.00 0.00 0.00	CTC	ACG TGC	ပ္ပ ဗ္ဗ	66 CC 45	CAC GTG	TAC	GTG	TGT	ပ္ပ ပ္ပ
AAG	ATG	AAG	ဗ္ဗ ဗ္ဗ	ပ္တပ္သ	TCA	GGA CCT	TTT	TCG	TGT	ပ္ပ ဗ္ဗ
TAG	ပ္ပ ဗ္ဗ ဗ္ဗ	AGG TCC	GTG	CTA	ည် ပို့ပို့	CAC	CT T	TAT ATA	ATG	ဗ္ဗ ပ္ပ ဗ္ဗ ပ္ဗ
CAG	CTG	AAC	GAC CTG	CCT	၁၁ ၁၁ ၁	GTG	TGC	ACG	TAT	CAA
99	ACG TGC	AAG	ဗ္ဗ ဗ္ဗ	၁၅၁	TAG	GAA	GAC	CGT	GAR	GCT CGA
S	<u>ပ</u> ္ပတ္သ	CTG	GTG	၁၅၅	GTC	AGT TCA	AAG TTC	TTT	TGT	C C
80	ATC	AAG	GAC	၁၅၁	ACC TGG	TTT	TAT ATA	CGA	TAA	S S S S S
	1486	1531	1576	1621	1666	1711	1756	1801	1846	1891

12/5€

1936 1981		GGC CCG ACT	ပ္ပတ္ပ	CGC		AAG CTG	AGT TCA TAT	ANA GAT	TAA TAA AAG	AGC ACT	AAA TTT AAA AAA TCG	ACG TGC	ACT TGA GAT	GAG CTC AAG	AAC
166 166	U U	TAC	TAA	ATT	ACA	GTA	ACG TGC						AGT	TAT	TAG
4 %	AGA	GTT	AGA	ATA TAT	TAA	GRA	GAA	GAT	CAA	TCT	ATT	<b>9</b> 000	TGA	AAA	GA A
<b>0 0</b>	GAT	AGT TCA	TAT ATA	GTT CAN	TAT	TTA	000 000 000 000	AAA	TTA	ACA	ATA TAT	ATA	දි දු	TGA	TTA
W 5-	ACC TGG	TTT	CGA	GTA	TAA	TAT	TGT	GAT	GAG	TCG	700 700	00 00 00 00 00	GTC	CAC	GTC
	ပ္ပ ပ္ပ	GTC	ACA	TGT	TTG	TTT AAA	CTG	ATG	CAC GTG	ACG TGC	TGA	CCN	၅ ၅ ၂၅	TTA AAT	TCG
C	767 <b>A</b> C <b>S</b>	AAG	ATG	GITT	CCA	10g	TCC	TGT ACA	000 000	၁၅ ၁၅ ၁၅	GAC	CCT GGA	CGA	CTA	AAT
2296	GAG	TAA	TTT	AAT	TTA AAT	TTG	CTG	TGA	TTA	CAT	TTT	AAT	GTG	TTG	ATT TAA
	ATC	TAC	CAT	AGG TCC	GTG	ATA	TAA	GTG	TGT	GAR	ATT	ACA TOT	ATA TAT	SE SE	AGT
2386	GTG	TCT	CGT	CGA	TAG	CIT	CCA	CAC	GAG	CAR	ည္ဟ	TIT	TGT	TTA	AGT

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CAC ACA GCA GCT ATC GAA GGT GTG CTC GTT CGG AAA ACA AAT TCA UO GAA CITI 2431 GAT TTA CTG ACA TGG ACA CTC GAC CCG CTA AAT GAC TGT ACC TGT GAG CTG GGC Fig.3 vi.

Total number of bases is: 2464.

Sequence ID 4.

ACTOGOGTGCTCTTCTCACCTGTTGCTCGGATTGTGTTGTACTAGAAAAA 30 20

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190 200 210 220 230 240 	4	250 260 270 280 290 300	v	310 320 330 340 350 360 360 360	<b>«</b>	370 380 390 400 410 420 	U
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230	<b>8</b>	290      TATA	<b>&gt;</b>	350  -  TCG1	Ø.	410  -  CCC	Δ,
2	×	ser 2	3	3 (S)	E	700	S
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0-8	•	g — ģ	<b>¤</b>	340 	阿	400    GCTC	H
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430 440 450 460 470 480 480 6 6 6 6 6 6 6 6 6 6 6 6 6 6 6 6 6 6 6	ı	;AAC	ø	Ž	Ŋ	BATK	Ω	X
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CAC	4	: SAGO	ø	i,	M	TGC.	ບ	GAG .
M	×	S S	Ω.	Ğ	S	299	O	GAA
460 	v	520    GCAG	ø	580  -  CAAA	×	640  -  -	Z	700 — GCAA
4 ( FAC)	>4	5; CAG	ø	200	O.	6 GTA	>	7
ACT	Ę+	CAG	ø	ACA	E٠	AGT	S	AGG
o — A	н	0 0 0 0	ρı	ACA ACA	E	0 — 1	Ś	0 0
450 	Ø	510 	a	570    CCGA	Ω,	630      GCTT	4	)  -    -    -    -  -  -  -  -  -  -  -
SATA TATA	Ω	Š	<u>α</u>	CCA	Ωı,	<b>Š</b>	ρ	<b>S</b> CA
S S	Ω	9		ATG	E	TCT	ល	
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430 GCTG	A	490 	E	550      CATG	X	610      TCGT	œ	670  -  GCGG
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			2210   	2220   GATCGCG
2230 2240 2250 2260 2270 2280 	2250   	2260      CTGAGTTGGT	2270   CACTCGGATA	2280     CGACTGT
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Fig.4 ix

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BMECR MSECR HVECR CLECR AGECR DMECR	BMECR Msecr Hvecr Ctecr Aaecr Dmecr	BMECR MSECR HVECR CLECR AAECR DMECR
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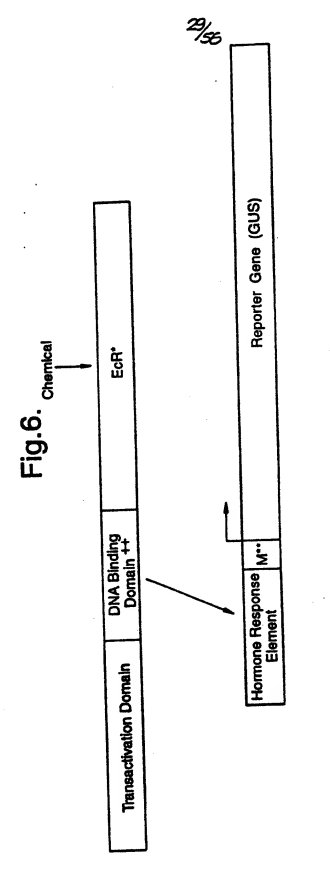
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ARROKKGPAPRQQEELCLVCGDRASGYHYNALTCEGCKGFFRRSVTKNAV PRRQKKGPAPRQQEELCLVCGDRASGYHYNALTCEGCKGFFRRSVTKNAV ARROKKGPAPRQQEELCLVCGDRASGYHYNALTCEGCKGFFRRSVTKNAV KSSSKKGPVPRQQEELCLVCGDRASGYHYNALTCEGCKGFFRRSVTKNAV AKKQKKGPTPRQQEELCLVCGDRASGYHYNALTCEGCKGFFRRSVTKNAV AKKSKKGPAPRVQEELCLVCGDRASGYHYNALTCEGCKGFFRRSVTKSAV	YICKFGHACEMDMYMRRKCQECRLKKCLAVGMRPECVIQEPS-KNKDRQR YICKFGHACEMDMYMRRKCQECRLKKCLAVGMRPECVVPESTCKNKRREK YICKFGHACEMDIYMRRKCQECRLKKCLAVGMRPECVVPENQCAMKRKEK YCCKFGHACEMDMYMRRKCQECRLKKCLAVGMRPECVVPENQCAIKRKEK YCCKFGHACEMDMYMRRKCQECRLKKCLAVGMRPECVVPENQCAIKRKEK YCCKFGRACEMDMYMRRKCQECRLKKCLAVGMRPGCVVPGNQCAMKREK	QKKDKGILLPVSTTTV	DPPPPEAARIHEVVPRYLSEKLMEQNRQKNIPPLSANQKSLIARL DPPPPEAARIHEVVPRFLTEKLMEQNRLKNVTPLSANQKSLIARL DPPPPEAARILECVQHEVVPRFLNEKLMEQNRLKNVPPLTANQKSLIARL DPPPHPMQQLLPEKLLMENRAKGTPQLTANQVAVIYKL DPPPHQAIPLLPEKLLQENRLRNIPLLTANQMAVIYKL BPPQHATIPLLPEKLLQENRLRNIPLLTANQMAVIYKL
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	BMECR MSECR HVECR CtECR AAECR DMECR	VEFAKGLPGFSKISQSDQITLLKASSSEVMMLRVARRYDAASDSVLFANN VEFAKGLPGFSKISQSDQITLLKASSSEVMMLRVARRYDAATDSVLFANN VEFAKGLPGFAKISQSDQITLLKACSSEVMMLRVARRYDAATDSVLFANN VEFAKGLPAFIKIPQEDQITLLKACSSEVMMLRMARRYDHDSDSILFANN VEFAKGLPAFTKIPQEDQITLLKACSSEVMMLRMARRYDHSSDSILFANN VEFAKGLPAFTKIPQEDQITLLKACSSEVMMLRMARRYDHSSDSIFFANN	459 401 418 384 524	
	BMECR MSECR HVECR CLECR ARECR DMECR	KAYTRDNYRQGGMAYVIEDLLHFCRCMFAMGMDNVHFALLTAIVIFSDRP QAYTRDNYRKAGMAYVIEDLLHFCRCMYSMSMDNVHYALLTAIVIFSDRP QAYTRDNYRKAGMAYVIEDLLHFCRCMYSMMDNVHYALLTAIVIFSDRP TAYTKQTYQLAGMEETIDDLLHFCRQMYALSIDNVETALLTAIVIFSDRP RSYTRDSYRMAGMADTIEDLLHFCRQMFSLTVDNVEYALLTAIVIFSDRP RSYTRDSYKMAGMADNIEDLLHFCRQMFSHTVDNVEYALLTAIVIFSDRP	509 451 434 574 574	136
	BMECR MSECR HVECR CLECR AAECR DMECR	GLEQPSLVEEIQRYYLNTLRIYIINQNSASSRCAVIYGRILSVLTELRTL GLEQPLLVEEIQRYYLKTLRVYILNQHSASPRCAVLFGKILGVLTELRTL GLEQPLLVEDIQRYYLNTLRVYILNQNSASPRGAVIFGEILGILTEIRTL GLEKAEMVDIIQSYYTETLKVYIVRDHGGESRCSVQFAKLLGILTELRTM GLEQAELVEHIQSYYIDTLRIYILNRHAGDPKCSVIFAKLLSILTELRTT GLEKAQLVEAIQSYYIDTLRIYILNRHCGDSMSLVFYAKLLSILTELRTL	559 501 518 518 542 624 624	

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GTQNSNMCISLKLKNRKLPPFLEEIWDVAEVARR	RNSSSSSSSSSSSNGSSNGNSSNSNSSQHGPHPHPHGQQLTPNQ RAERMRASVGGAITAGIDCDSASTSAAAAAAAQHQPQPQPQPQPQPSSLTQND		HANGSGSGGSNNNSSSG
BMBCR MSECR HVECR CLECR AAECR DMECR	BMECR MSECR HVECR CLECR AAECR DMECR	BMECR MSECR HVECR CLECR ABECR DMECR	BMECR MSECR HVECR CLECR ARECR

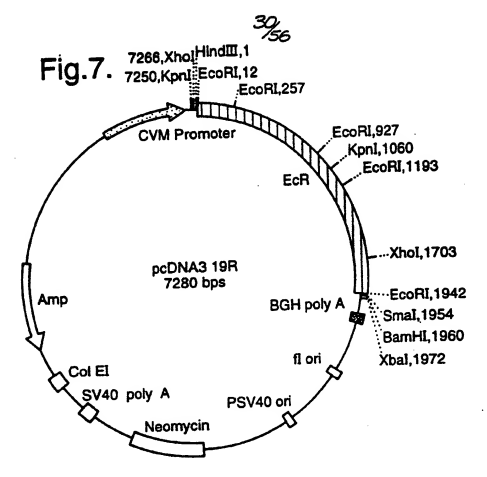
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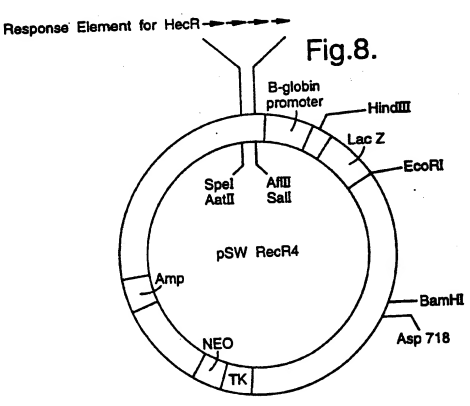
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A B B C R D M B C R	PMGNG	PMGNGVGVGVGVGGNVSMYANAQTAMALMGVALHSHQQQLIGGVAVKSEH
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AAECR	1 1	675
Target B	Citation	878



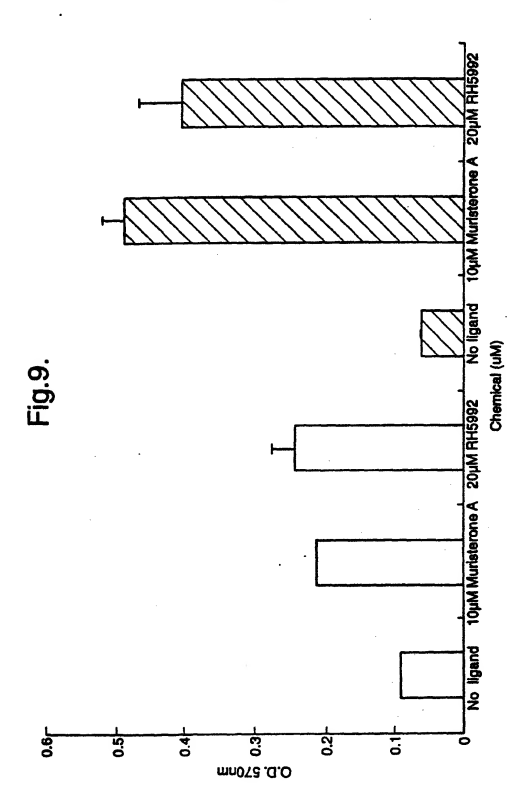
++ Glucocorticoid receptor DNA binding and transactivation domains
• Insect ecdysone ligand binding domain
•• Minimal 35S CaMV promoter

PCT/GB96/01195





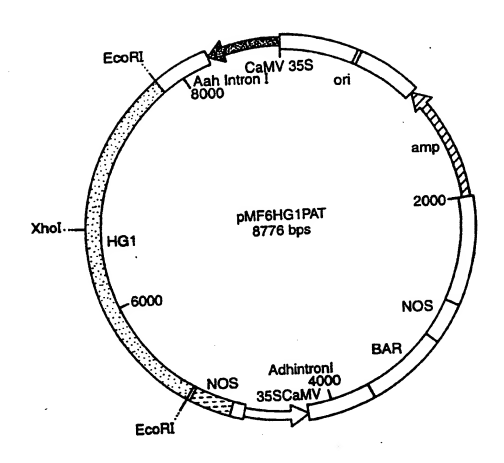
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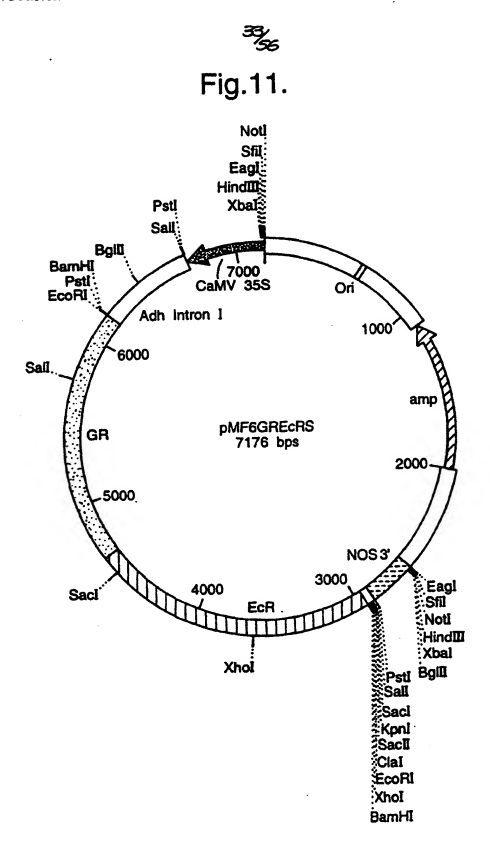


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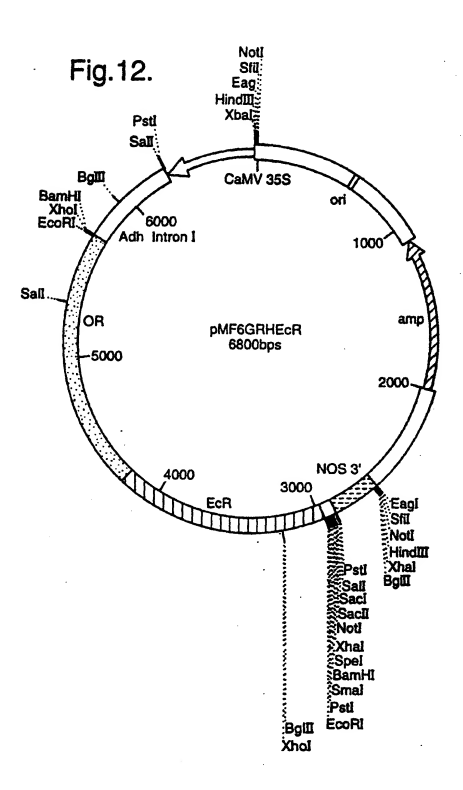
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Fig.10.

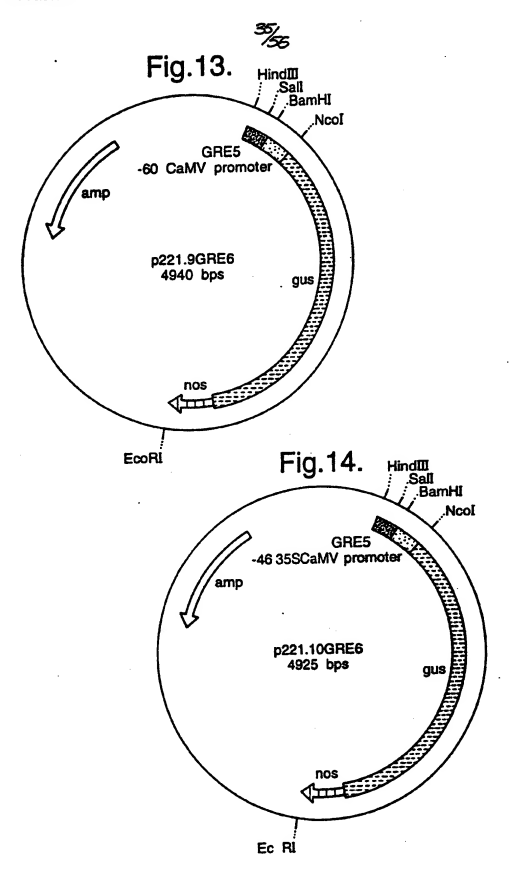




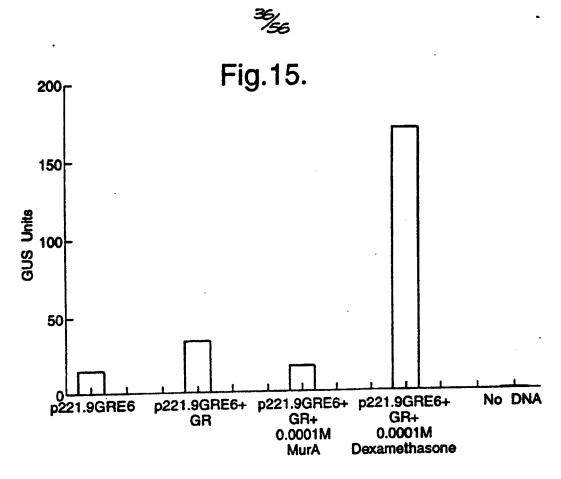


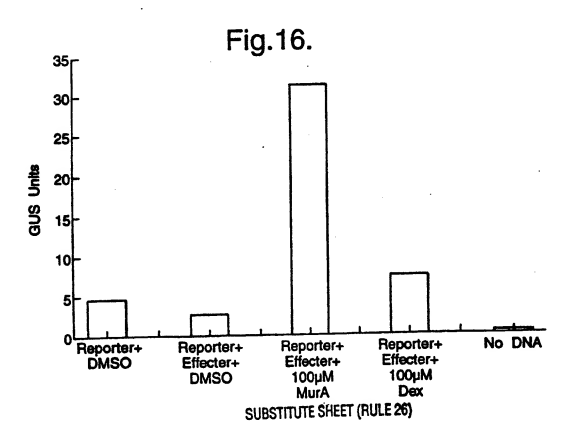


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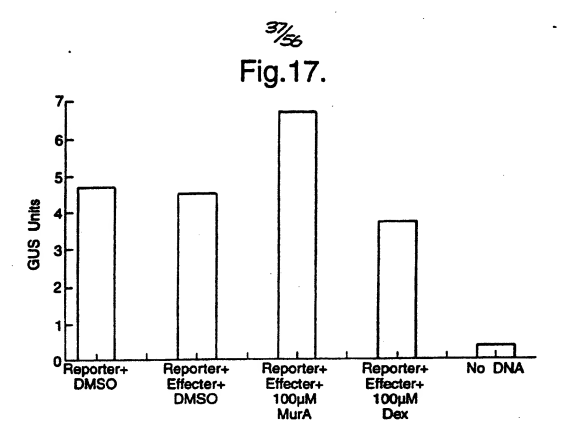


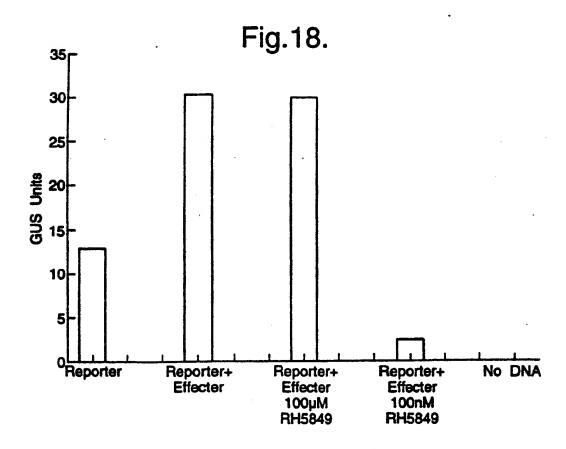
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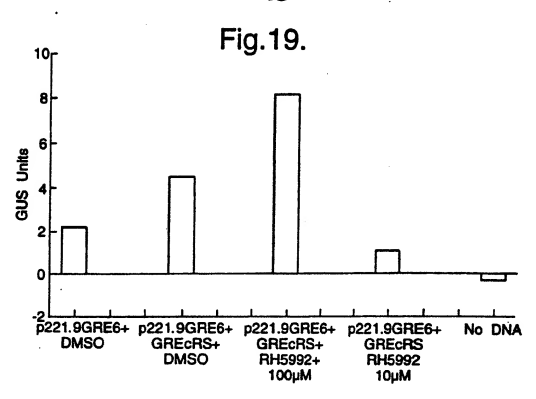
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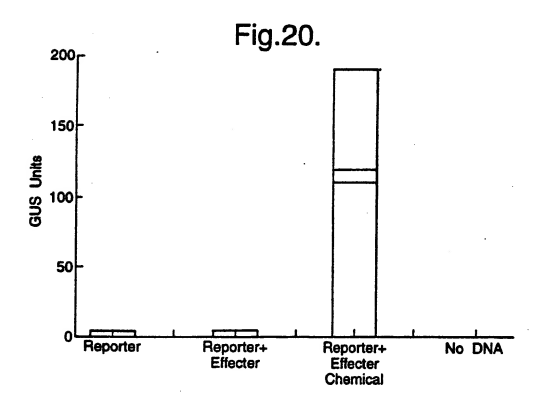




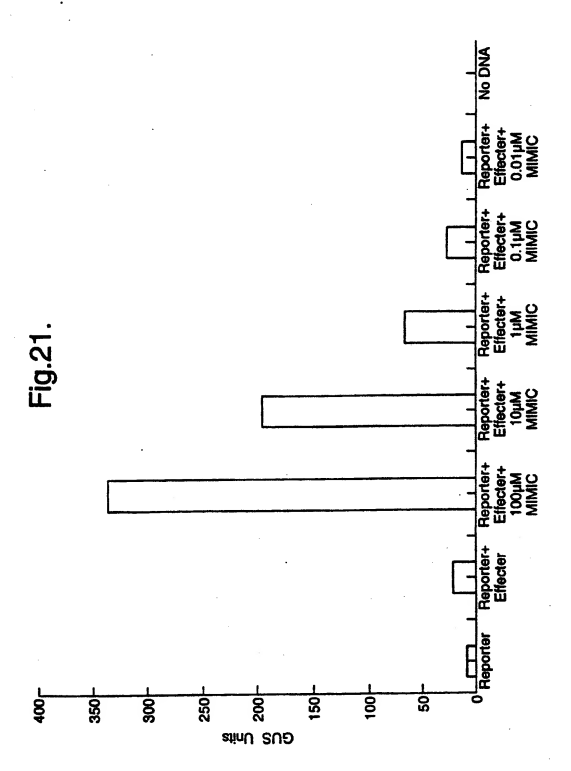
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Fig.22. NotI Eagl HindIII Pstl EcoRl 35SCam V ori Sall `6000 1000 GR Amp R pMF7GREcRS 6700 bps -5000 2000\_ 4000 3000 NOS HEcR Eagl Sfil Xhoi Bgtti Noti HindⅢ Xbai ĖgⅢ BamHI



Fig.23. Noti Sfil Eagl Hind Xbal Psti EcoRI: 35SCamV ori Sall **6000** 1000 GR -5000 pMF7GRHEcR 6500 bps Amp R 2000 4000 3000 NOS Eagl Shi Noti Xhol: BgIII BamHİ ĴHindⅢ Xbal BgⅢ

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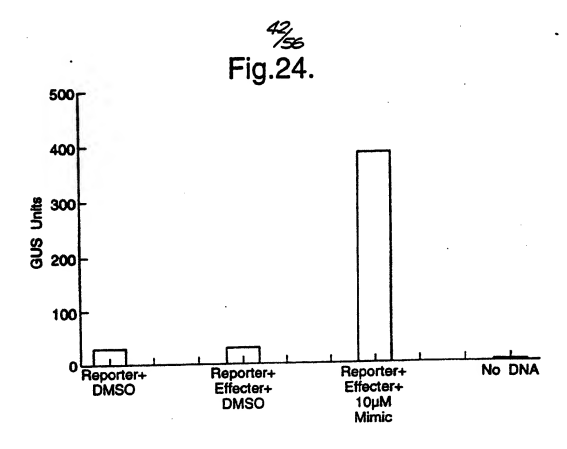
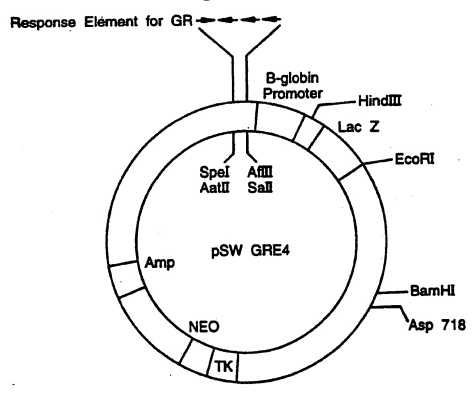


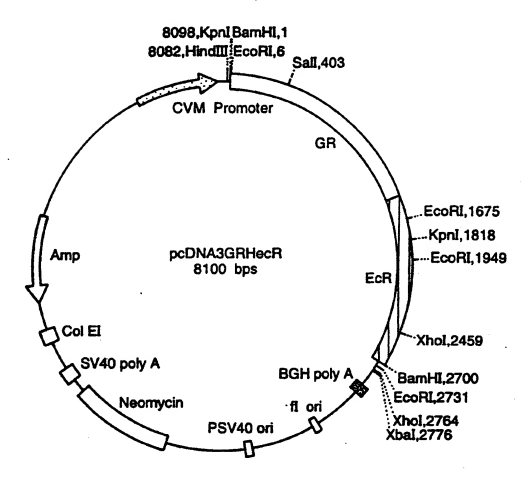
Fig.26.



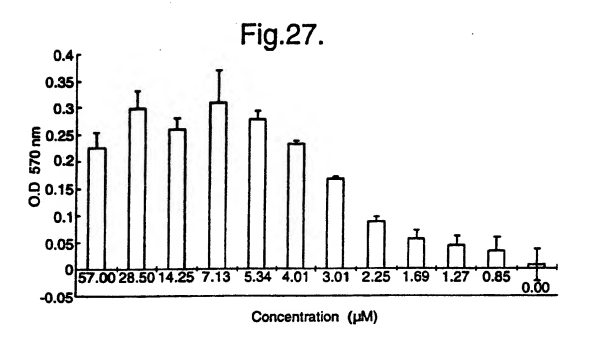
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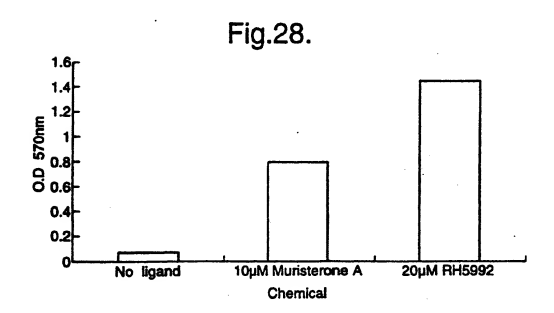


Fig.25.

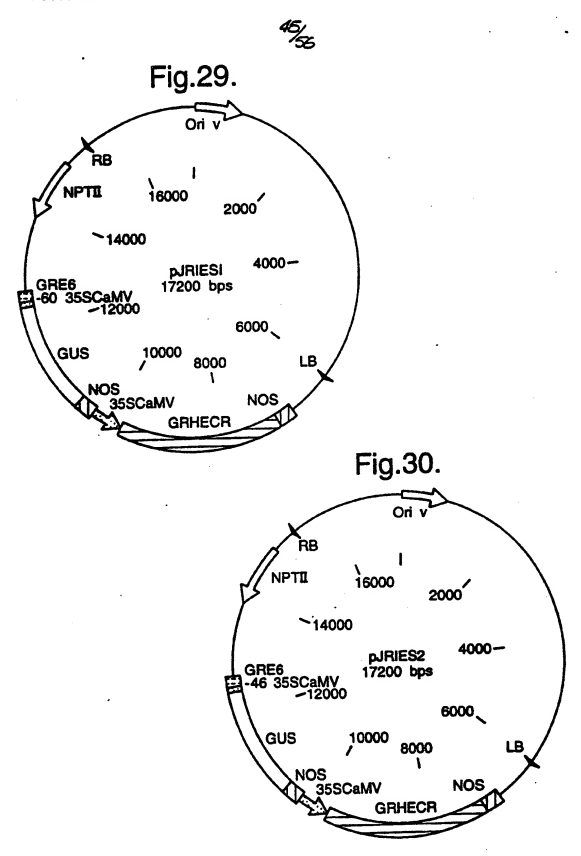


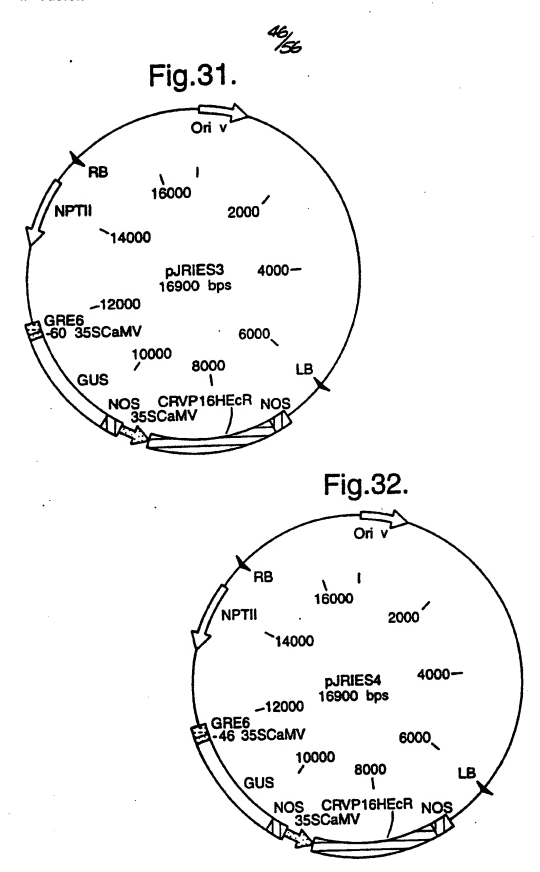
44,56





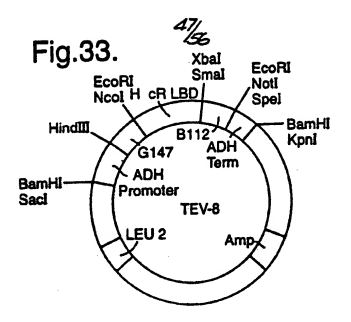
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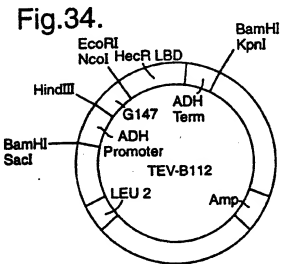


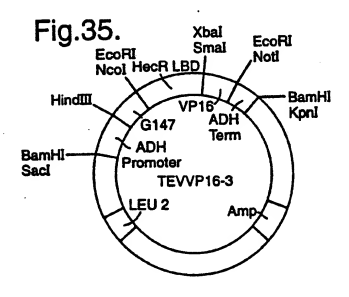


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48, 55

Fig.36.

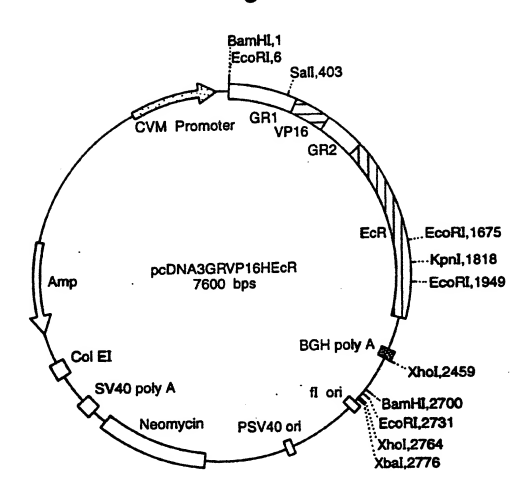
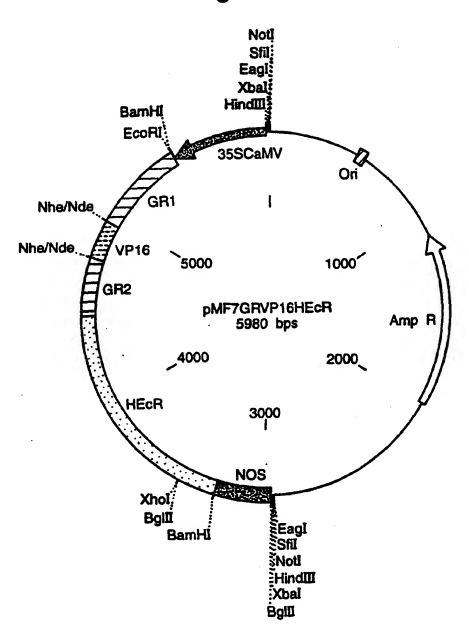


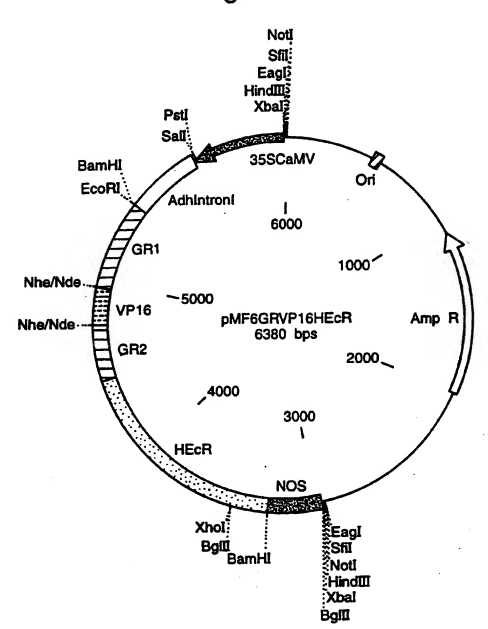


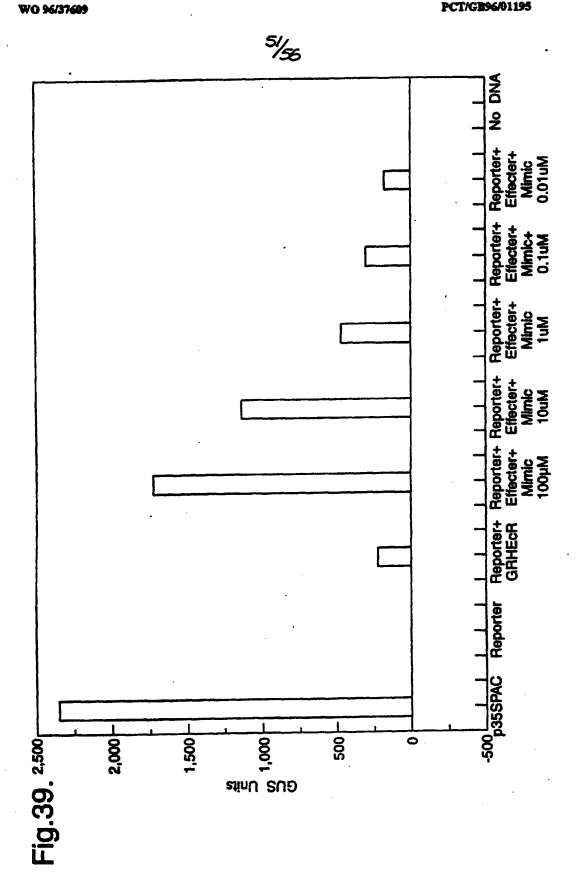
Fig.37.



50<sub>55</sub>

Fig.38.





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CTG

**E S** 

SCT CGA

GAG

36c 3cg

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**366** 

Ag to

**4**00

GAT

271

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Spodoptera exigna DNA sequence.

Fig.40.

Sequence ID (

SPODOPTERA EXIGUA HINGE AND LIGAND BINDING DOMAINS

45	AGG	AGT	GAT	CGA	GTG
	AAA TTT	GTC	TGT	CCA	AAT
39	ATG	CCA TOO	CAG	GTG	AAG
	GC T	TTG	ATG	OTO CAC	CTC
33	TGT ACA	AAG	ATT	CIC	AGG
	CAG GTC	GAC	သည် ၁၃၃	CAC	ACA
27	AAC	AAA	CCT	att	AGG TCC
	GAA	GAA	ATG	AGA	GAC
21	CCA GGT	AGG TCC	CAC GTG	GCA	atg
	GTG	ST	GAT	ပ္ပ ပ္ပ ပ္ပ	CTA
15	GTG	GCA CGT	GAT	GAG	AAG
	TGC	AAG	GTG	CCA	GAA
o –	GAG	AAT T	ACA	GGA	AAT
	ပ္ပ ပ္ပ	GAG	ACG TGC	ပ္ပဲ ပိပ္ပဲ ပိ	CTG
m —	100 100	ANA	ACA	CCA	TTC
	<b>ન</b>	46	91	136	181

GAT

80

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**6** 600

CTG GAC

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88
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GAC
5 <b>3</b>
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हु हु
GAT
TAA
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GATA TAB
811

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SE
CG A
SE
ACT
GAG
GTA
CAC CAC
CAT
GTT
CTC
CTT
යි දුර
CAT GTA
856

946 AAA TTT

Total number of bases is: 948.

55/5E

Fig.41

Sequence I.D. 7

SECR Teg clone comparison between Heliothis 19R clone and Sequence

RPECVVPENQCAMKRKEKKAQREKDKLPVSTTTVDDHMPPIMQCDPPPPEAARILECVQ RPECVVPENOCAMKRKEKKAOREKDKLPVSTTTVDDHMPPIMOCDPPPPEAARI HECR SECR

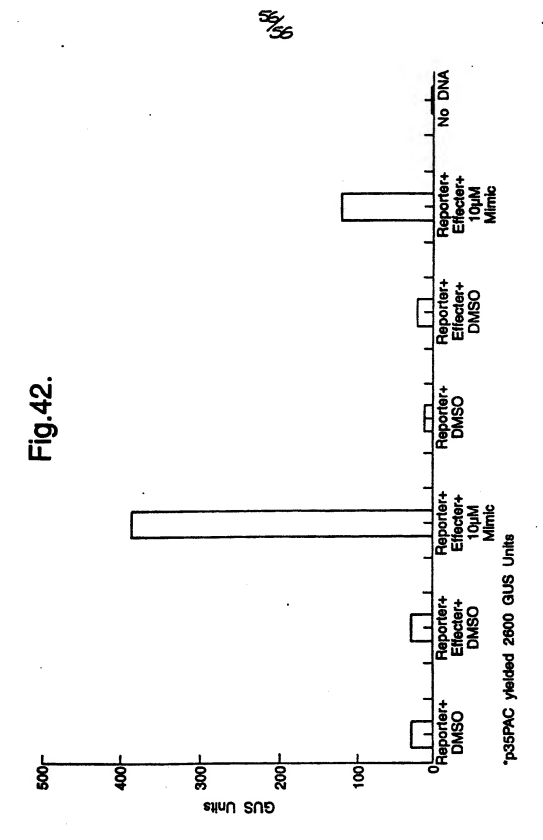
**HEVVPRFLNEKLMERTRLRNVPPLTANQKSLIARLVWYQEGYEQPSEEDLKRVTQSD** HEVVPRFLNEKLMEGNRLKNVPPLTANOKSLIARLVWYQEGYEQPSEEDLKRVTQSD HECR SECR

**EDDEDSDMPFRQITEMTILTVQLIVEFAKGLPGFAKISQSDQITLLKACSSEVMMLR** edeeesdmpfroitemtiltvolivefakglpafakisqsdqitllkacssevmlr HECR SECR

**VARRYDAATDSVLFANNOAYTRDNYRKAGMAYVTEDLLHFCRCMYSMMDNVHYALL VARRYDAATDSVLFANNQAYTRDNYRKAGMAYVIEDLLHFCRCMYSMMDNVHYALL** HECR SECR

TAIVIFSDRPGLELTLLVEELQRYYLNTLRVYILNQNSRSPCCPVIYAKILGILTEL Taivifsdrpgleopllveeiorytlntlrvyilnonsasprgavifgeilgiltei HECR SECR

HECR RTLOMONSNMCISLKLKKRKLPPFLEEIDWDV SECR RTLOMONSNMCISLKLKNRNVPPFFEDIDWDV



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mail Application No

PCT/GB 96/01195 A. CLASSIFICATION OF SUBJECT MATTER 1PC 6 C12N15/12 C12N15/85 C07K14/72 C07K19/00 C12N15/62 A61K38/16 C12N5/10 According to International Patent Classification (IPC) or to both national classification and IPC B. PIELDS SEARCHED m documentation searched (classification system followed by classification symbols) CO7K C12N A01N IPC 6 Documentation searched other than minimum documentation to the extent that such documents are included in the fields searched Electronic data base exempted during the international search (name of data base and, where practical, search terms used) C. DOCUMENTS CONSIDERED TO BE RELEVANT Referent to claim No. Citation of document, with indication, where appropriate, of the relevant passages 4,5,44, MO,A,93 83162 (GENENTECH INC) 18 February X 92-99 1,3, Y see abstract: claims 1-27: figure 1 8-43, 45-49, 51-91 4,5,44, X WO.A.91 13167 (UNIV LELAND STANFORD 50,93-99 JUNIOR) 5 September 1991 2.3 Y see abstract; claims 2,24 -/--Patent family members are listed in annex. Purther documents are listed in the continuation of box C. cial categories of cited documents: "I" later document published after the international filing door priority date and not in conflict with the application cited to understand the principle or theory underlying the invention. connect defining the general state of the set which is not anishmed to be of perticular relevances tier document but published on or after the interestional "X" document of particular relevance; the claimed in cannot be considered novel or cannot be consid-involve an inventive step when the document is document which may throw doubts on priority claim(s) or which is cleal to establish the publication date of another classion or other special reason (as specified) document of particular reference; the daimed invention cannot be considered to insulve an inventive step when the document is combined with one or more other such documents, such combination being obvious to a person skilled in the set. "O" document referring to an oral dischoose, use, exhibition or current published polar to the inte ter than the polarity date claimed "A" document member of the same patent family Date of mailing of the international search report Date of the actual exempletion of the international search 19.08.96 9 August 1996 Authorized officer Name and mailing address of the BA European Painnt Office, P.R. 5818 Patentiann 2 NL - 2250 HV Rijevijk Tul. (+31-70) 340-2040, Tz. 34 651 epo nl. Parc (+31-70) 340-3016

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	Ohervations where certain claims were found unsearchable (Continuation of item 1 of first sheet)
This int	ernational bearch report has not been established in respect of certain claims under Article 17(2)(a) for the following reasons:
ı. <b>X</b>	Claims Nos.: 98 because they relate to subject matter not required to be searched by this Authority, mannely: Although this claim is directed partly to a method of treatment of the human/animal body the search has been carried out and based on the alleged effects of the compound/composition
² []	Claims Nos.: because they relate to parts of the international application that do not comply with the prescribed requirements to such an extent that no meaningful international search can be carried out, specifically:
3.	Claims Nos.: because they are dependent claims and are not drafted in accordance with the second and third sentences of Rule 6.4(a).
Bex II	Observations where unity of invention is lacking (Continuation of item 2 of first sheet)
This Inte	ernational Searching Authority found multiple inventions in this international application, as follows:
1.	As all required additional search fees were timely paid by the applicant, this international search report covers all searchable claims.
2 🔲	As all searchable claims could be searches without effort justifying an additional fee, this Authority did not invite payment of any additional fee.
3. 🔲 j	As only some of the required additional search fees were timely paid by the applicant, this international search report covers only those claims for which fees were paid, specifically claims Nos.:
	No required additional search fees were timely paid by the applicant. Consequently, this international search report is restricted to the investion first mentioned in the claims; it is covered by claims Nos.:
<b>.</b>	The additional search fees were accompanied by the applicant's protest.

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